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(12)

EUROPEAN PATENT SPECIFICATION

- (45) Date of publication and mention of the grant of the patent: 14.08.1996 Bulletin 1996/33
- (21) Application number: 92810056.9
- (22) Date of filing: 27.01.1992

- (51) Int CL⁶: **C12N 15/62**, C12N 1/21, G01N 33/574, A61K 39/395, C07K 16/30
 // (C12N1/21, C12R1:19)
- (54) Recombinant antibodies specific for a growth factor receptor Rekombinante Antikörper spezifisch für einen Wachstumsfaktor-Rezeptor

Rekombinante Antikörper spezifisch für einen Wachstumsfaktor-Rezeptor

Anticorps recombinants spécifiques pour un récepteur de facteur de croissance

- (84) Designated Contracting States:
 AT BE CH DE DK ES FR GB GR IT LI LU NL PT SE
- (30) Priority: 05.02.1991 EP 91810079
- (43) Date of publication of application: 09.09.1992 Bulletin 1992/37
- (83) Declaration under Rule 28(4) EPC (expert solution)
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EP 0 502 812 B1

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Description

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Background of the invention

Growth factors and their receptors are involved in the regulation of cell proliferation, and they also seem to play a role in tumor growth. The c-erbB-2 growth factor receptor protein, a protein of the membrane receptor protein tyrosine kinase family (A. Ullrich & J. Schlessinger, Cell 61: 203-212, 1990), is found in human breast tumors and human ovarian carcinomas. Amplification of the c-erbB-2 gene and over-expression of the protein appears to correlate with poor prognosis for tumor patients. Thus the c-erbB-2 protein has potential, both as a diagnostic marker and as a target for cancer therapy. Sequence analysis reveals that c-erbB-2, also called HER2, a glycoprotein of 185 kilo-Dalton (gp185), is identical or closely related to the human analog of the neu oncogene (A.L. Schechter et al., Science 229: 976-978, 1985) and shows considerable sequence homology to the human epidermal growth factor (EGF) receptor.

Of particular interest in tumor diagnosis and therapy are antibodies directed to tumor markers. Polyclonal antibodies may be obtained from the serum of mammals immunized with the antigen, i.e. the tumor marker. The development of hybridoma technology made it possible to generate continuous cell lines, in particular murine hybridomas, producing monoclonal antibodies of the desired specificity. Murine monoclonal antibodies directed to c-erbB-2 are known and are described, for example, by S.J. McKenzie et al., Oncogene 4: 543-548, 1989; R.M. Hudziak et al., Molecular and Cellular Biology 9: 1165-1172, 1989; International Patent Application WO 89/06692 (Genentech); and Japanese Patent Application Kokai 02-150 293 (Ajinomoto KK).

A major limitation in the use of murine-derived monoclonal antibodies as <u>in vivo</u> diagnostic and therapeutic agents is their immunogenicity as foreign proteins, their rather long persistence in the circulation, and the formation of damaging immune complexes. On the other hand, the treatment with human monoclonal antibodies is also limited since human hybridoma cell lines are hard to prepare, generally unstable, and do not produce monoclonal antibodies of appropriate specificity in sufficient quantities and at reasonable costs. In principle, the <u>in vitro</u> use of murine monoclonal antibodies is without limitation. However, production costs of monoclonal antibodies and, depending on the type of immunoassay used, the need for attaching a detectable marker to the antibody make it desirable to find more economic alternatives to regular murine monoclonal antibodies.

A promising alternative is the modification of immunoglobulin genes in order to tailor antibodies for particular diagnostic and therapeutic tasks. Due to the fact that the variable region and each of the constant region domains of immunoglobulin molecules are encoded in separate exons with their own splice sites, recombinant DNA techniques can be used to isolate different parts of cloned immunoglobulin genes and ligate them to parts of other immunoglobulins or to effector molecules. The reconstructed genes are expressed by appropriate transformed continuous cell lines. Murine antibodies can, for example, be converted into "humanized" antibodies by exchanging murine constant domain exons for human immunoglobulin constant domain exons, thus generating chimeric antibodies with murine antibodycombining sites and human constant domains. The chimeric antibodies retain the antigen specificity determined by the murine variable domains, but also exhibit human effector functions (such as complement binding, stimulation of phagocytosis, triggering of granule release by mast cells) determined by the carboxy-terminal constant domain segments of the heavy chain polypeptides. An even more sophisticated technique in tailoring antibodies described in European Patent Application 0 239 400 exchanges also other fairly conserved domains, the so-called framework regions (FRs), within the murine variable domains for corresponding framework regions from human antibodies or for other human protein sequences. Such an antibody should be even less immunogenic in man since the only parts derived from a murine antibody are those hypervariable regions which define a particular specificity for an antigen, the so-called complementarity determining regions (CDRs).

Furthermore, fusion proteins different from immunoglobulins may be formed, e.g. single-chain antibodies, which retain the specificity and binding properties of the starting murine monoclonal antibody, but have otherwise novel properties derived from the non-immunoglobulin part of the fusion protein. The smallest domain of a monoclonal antibody which can bind to the antigen is the so-called Fv fragment which consists of the variable domains of the heavy and light chains. Fv fragments are difficult to prepare by proteolytic techniques since the corresponding variable domains tend to dissociate upon dilution. Fv molecules constructed by joining the variable domains of the heavy and light chains via a short peptide linker, also called single-chain antigen binding proteins, bind to an antigen with similar characteristics as the original monoclonal antibody (R.E. Bird et al., Science 242: 423-426, 1988; J.S.Huston et al., Proc. Natl. Acad. Sci. USA 85: 5879-5883, 1988; and International Patent Application WO 89/09825 (Celltech)). Fv encoding genes can, in principle, be linked to genes encoding effector molecules by recombinant gene technology. It is known, for example, that Fv encoding gene sequences can be linked to a gene encoding a portion of the Pseudomonas exotoxin A gene (V.K. Chaudhary et al., Nature 339: 394-397, 1989; and International Patent Application WO 89/11533 (I. Pastan et al.)).

Object of the invention

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It is an object of this invention to provide recombinant antibodies directed to the extracellular domain of the human growth factor receptor c-erbB-2 comprising a light chain variable domain and a heavy chain variable domain of a monoclonal antibody, monoclonal antibodies directed to c-erbB-2 themselv s, a method of manufacture of said recombinant antibodies and said monoclonal antibodies, hybridoma cells secreting said monoclonal antibodies, a method of manufacture of said hybridoma cells, DNA coding for the heavy chain variable domain, for the light chain variable domain and for the recombinant antibody, a method of manufacture of said DNA, hybrid vectors suitable for expression of said DNA, host cells transformed with said DNA, and the use of said recombinant antibodies and said monoclonal antibodies in the diagnosis and treatment of tumors.

Detailed description of the invention

The invention concerns a recombinant antibody directed to the extracellular domain of the growth factor receptor c-erbB-2, a human glycoprotein of 185 kilo-Dalton (gp185), comprising a heavy chain variable domain and a light chain variable domain of a monoclonal antibody.

Such a recombinant antibody may be a chimeric antibody consisting, for example, of a mouse heavy chain variable domain with the specificity for c-erbB-2 and a human heavy chain constant domain α , γ , δ , ϵ , or μ , preferably γ , such as γ 1 or γ 4, and of a mouse light chain variable domain with the specificity for c-erbB-2 and a human light chain constant domain κ or λ , preferably κ , all assembled to give a functional antibody.

The preferred recombinant antibody of the invention is a single-chain antibody wherein the heavy chain viable domain and the light chain variable domain are linked by way of a spacer group, preferably a peptide. Most preferred is a single-chain antibody wherein the heavy chain variable domain is located at the N-terminus of the recombinant antibody. The single-chain recombinant antibody may further comprise an effector molecule and/or signal sequences facilitating the processing of the antibody by the host cell in which it is prepared. Effector molecules considered are those useful for diagnostic or therapeutic purposes, for example enzymes causing a detectable reaction, e.g. phosphatase, such as alkaline phosphatase from <u>E.coli</u> or mamalian alkaline phosphatase, e.g. bovine alkaline phosphatase, horseradish peroxidase, β-D-galactosidase, glucose oxidase, glucoamylase, carbonic anhydrase, acetylcholinesterase, lysozyme, malate dehydrogenase or glucose-6-phosphate, a peptide having particular binding properties, e.g. streptavidin from <u>Streptomyces avidinii</u> strongly binding to biotin, or enzymes, toxins or other drugs attacking the cells to which the antibody is bound, e.g. a protease, a cytolysin or an exotoxin, for example ricin A, diphtheria toxin A, or <u>Pseudomonas</u> exotoxin. In the following a single-chain recombinant antibody further comprising an effector molecule is referred to as fusion protein or intended to be within the meaning of the terms "single chain (recombinant) antibody" or "recombinant antibody", if appropriate.

The term effector molecule also includes biologically active variants of the above-mentioned proteins, e.g. variants produced from a DNA which has been subjected to in vitro mutagenesis, with the provision that the protein encoded by said DNA retains the biological activity of the native protein. Such modifications may consist in an addition, exchange or deletion of amino acids, the latter resulting in shortened variants. For example, an enzyme, such as phosphatase, may be prepared from a DNA which has been modified to facilitate the cloning of the encoding gene, or an exotoxin, such as Pseudomonas exotoxin, may be prepared from a DNA which has been mutated to delete the cell binding domain.

The recombinant antibodies of the invention are tested for their specificity to the extracellular domain of c-erbB-2, for example by immunofluorescent staining of cells expressing high levels of c-erbB-2, by immunoblotting either directly or by way of immunoprecipitation and protein blotting of the immunocomplexes, or by another immunoassay such as a binding, crossinhibition or competition radio- or enzyme immunoassay.

The variable domain of an antibody heavy or light chain consists of so-called framework regions (FRs), which are fairly conserved in antibodies with different specificities, and of hypervariable regions also called complementarity determining regions (CDRs), which are typical for a particular specificity.

Preferred recombinant antibodies of the invention are those wherein the heavy chain variable domain comprises a polypeptide of the formula

$$FR_1 - CDR_{1H} - FR_2 - CDR_{2H} - FR_3 - CDR_{3H} - FR_4$$
 (I)

wherein FR₁ is a polypeptide residue comprising at least 25-29, preferably 25-33 naturally occurring amino acids, FR₂ is a polypeptide residue comprising 12-16 naturally occurring amino acids, FR₃ is a polypeptide residue comprising 30-34 naturally occurring amino acids, FR₄ is a polypeptide residue comprising at least 6-10, preferably 6-13 naturally occurring amino acids, CDR_{1H} is a polypeptide residue of the amino acid sequence 32 to 36 of SEQ ID NO:4, CDR_{2H} is a polypeptide residue of SEQ ID NO:4, and CDR_{3H} is a polypeptide residue of

the amino acid sequence 100 to 109 of SEQ ID NO:4, or, CDR_{1H} is a polypeptide residue of the amino acid sequence 32 to 36 of SEQ ID NO:8, CDR_{2H} is a polypeptide residue of the amino acid sequence 51 to 67 of SEQ ID NO:8, and CDR_{3H} is a polypeptide residue of the amino acid sequence 100 to 110 of SEQ ID NO:8, and wherein the amino acid Cys may be in the oxidized state forming S-S-bridges. These particular complementarity determining regions are Asn-Tyr-Gly-Met-Asn (CDR_{1H}), Trp-IIe-Asn-Thr-Ser-Thr-Gly-Glu-Ser-Thr-Phe-Ala-Asp-Asp-Phe-Lys-Gly (CDR_{2H}), and Trp-Glu-Val-Tyr-His-Gly-Tyr-Val-Pro-Tyr (CDR_{3H}) according to SEQ. ID NO: 4, or Ser-Tyr-Trp-Met-Asn (CDR_{1H}), Met-IIIe-Asp-Pro-Ser-Asp-Ser-Glu-Thr-Gln-Tyr-Asn-Gln-Met-Phe-Lys-Asp (CDR_{2H}) and Gly-Gly-Ala-Ser-Gly-Asp-Trp-Tyr-Phe-Asp-Val (CDR_{3H}) according to SEQ. ID NO:8.

Especially preferred are recombinant antibodies comprising a heavy chain variable domain of formula I, wherein the polypeptide residues of the framework regions FR₁, FR₂, FR₃ and FR₄ are those preferably occurring in mammalian, especially murine or human, antibodies.

In a first embodiment of the invention, most preferred are recombinant antibodies with a heavy chain variable domain comprising a polypeptide of the amino acid sequence 2 to 120, of SEQ ID NO:4, wherein optionally one or more, e.g. 1, 2, 3 or 4, single amino acids within the amino acid sequences 2 to 31 (FR₁), 37 to 50 (FR₂), 68 to 99 (FR₃), and/or 110 to 120 (FR₄), are replaced by other amino acids or deleted, and wherein the amino acid Cys may be in the oxidized state forming S-S-bridges, in particular the recombinant antibodies with a heavy chain variable domain comprising a polypeptide of the amino acid sequence 2 to 120 of SEQ ID NO:4, wherein the amino acid Cys may be in the oxidized state forming S-S-bridges.

In a second embodiment of the invention, most preferred are recombinant antibodies wherein the heavy chain variable domain comprises a polypeptide of the amino acid sequence 2 to 121, of SEQ ID NO: 8, wherein optionally one or more, e.g. 1, 2, 3 or 4, single amino acids within the amino acid sequences 2 to 31 (FR₁), 37 to 50 (FR₂), 68 to 99 (FR₃), and/or 111 to 121 (FR₄), are replaced by other amino acids or deleted, and wherein the amino acid Cys may be in the oxidized state forming S-S-bridges, in particular the recombinant antibodies with a heavy chain variable domain comprising a polypeptide of the amino acid sequence 2 to 121 of SEQ ID NO: 8, wherein the amino acid Cys may be in the oxidized state forming S-S-bridges.

For example, a hydrophobic amino acid within the framework regions may be replaced by another amino acid, preferably also a hydrophobic amino acid, e.g. a homologous amino acid, replaced by two amino acids, or deleted. Likewise, a hydrophilic amino acid within the framework region may be replaced by another amino acid, two amino acids or deleted, whereby replacing amino acids preferably maintain the hydrogen bond structure of the corresponding framework region.

Likewise preferred recombinant antibodies of the invention are those wherein the light chain variable domain comprises a polypeptide of the formula

$$FR_6 - CDR_{11} - FR_7 - CDR_{21} - FR_8 - CDR_{31} - FR_9$$
 (II)

wherein FR₆ is a polypeptide residue comprising naturally occurring amino acids, preferably 19-25, especially 19-23 naturally occurring amino acids, FR₇ is a polypeptide residue comprising 30-34 naturally occurring amino acids, FR₈ is a polypeptide residue comprising 30-34 naturally occurring amino acids, FR₉ is a polypeptide residue comprising naturally occurring amino acids, particularly 7-11 naturally occurring amino acids, and CDR_{1L} is a polypeptide residue of the amino acid sequence 159 to 169 of SEQ ID NO:4, CDR_{2L} is a polypeptide residue of the amino acid sequence 185 to 191 of SEQ ID NO:4, and CDR_{3L} is a polypeptide residue of the amino acid sequence 224 to 232 of SEQ ID NO:4, or CDR_{1L} is a polypeptide residue of the amino acid sequence 160 to 170 of SEQ ID NO:8, CDR_{2L} is a polypeptide residue of the amino acid sequence 186 to 192 of SEQ ID NO:8, and CDR_{3L} is a polypeptide residue of the amino acid sequence 225 to 232 of SEQ ID NO:8, and wherein the amino acid Cys may be in the oxidized state forming S-S-bridges. These particular complementarity determining regions are Lys-Ala-Ser-Gln-Asp-Val-Tyr-Asn-Ala-Val-Ala (CDR_{1L}), Ser-Ala-Ser-Ser-Arg-Tyr-Thr (CDR_{2L}), and Gln-Gln-His-Phe-Arg-Thr-Pro-Phe-Thr (CDR_{3L}) according to SEQ ID No:8, or Lys-Ala-Ser-Gln-Asp-Ile-Lys-Lys-Tyr-Ile-Ala (CDR_{1L}), Tyr-Thr-Ser-Val-Leu-Gln-Pro (CDR_{2L}) and Leu-His-Tyr-Asp-Tyr-Leu-Tyr-Thr (CDR_{3L}) according to SEQ ID No. 8.

Especially preferred are recombinant antibodies comprising a light chain variable domain of formula II, wherein the polypeptide residues of the framework regions FR₅, FR₆, FR₇ and FR₈ are those preferably occurring in mammalian, especially murine or human, antibodies.

In one embodiment of the invention, most preferred are recombinant antibodies wherein the light chain variable domain comprises a polypeptide of the amino acid sequence 136 to 241 of SEQ ID NO:4, wherein optionally one or more, e.g. 1, 2, 3 or 4, single amino acids within the amino acid sequences 136 to 158 (FR₆), 170 to 184 (FR₇), 192 to 223 (FR₈), and/or 233 to 241 (FR₉) are replaced by other amino acids or deleted, and wherein the amino acid Cys may be in the oxidized state forming S-S-bridges, in particular the recombinant antibodies with a light chain variable domain comprising a polypeptide of the amino acid sequence 136 to 241 of SEQ ID NO:4, wherein the amino acid Cys may be in the oxidized state forming S-S-bridges.

In a second embodiment of the invention, most preferred are recombinant antibodies wherein the light chain variable domain comprises a polypeptide of the amino acid sequence 137 to 241 of SEQ ID NO:8, wherein optionally one or more, e.g. 1, 2, 3 or 4 single amino acids within the amino acid sequences 137 to 159 (FR₆), 171 to 185 (FR₇), 193 t 224 (FR₈), and/or 233 to 241 (FR₉) are replaced by other amino acids or deleted, and wherein the amino acid Cys may be in the oxidized state forming S-S-bridges, in particular the recombinant antibody wherein the light chain variable domain comprises a polypeptide of the amino acid sequence 137 to 241 of SEQ ID NO:8, wherein the amino acid Cys may be in the oxidized state forming S-S-bridges.

For example, amino acids within the framework regions may be replaced by other amino acids or deleted as detailed above for the heavy chain.

Especially preferred is a single-chain recombinant antibody wherein the heavy chain variable domain and the light chain variable domain are linked by way of a spacer group consisting of 10 to 30, e.g. around 15, amino acids, in particular a single-chain recombinant antibody comprising a polypeptide of the formula

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$$\mathsf{FR_1\text{-}CDR_{1H}\text{-}FR_2\text{-}CDR_{2H}\text{-}FR_3\text{-}CDR_{3H}\text{-}FR_4\text{-}Sp\text{-}FR_6\text{-}CDR_{1L}\text{-}FR_7\text{-}CDR_{2L}\text{-}FR_8\text{-}CDR_{3L}\text{-}FR_9}} \tag{III),}$$

wherein FR₁,CDR_{1H},FR₂,CDR_{2H},FR₃,CDR_{3H},FR₄,FR₆,CDR_{1L},FR₇,CDR_{2L}, FR₈,CDR_{3L} and FR₉ have the meanings as mentioned before and Sp is a peptide spacer consisting of about 10 to 30, e.g. around 15, amino acids; and wherein the heavy chain or the light chain variable domain is further connected to an effector molecule, e.g. an enzyme, such as phosphatase, particularly alkaline phosphatase, or a toxin, such as Pseudomonas exotoxin, or a variant thereof. Preferably, the effector molecule is connected to the light chain variable domain, optionally via a peptide spacer consisting of one or more, e.g. 1-10 amino acids.

These fusion proteins comprising a single-chain recombinant antibody and an effector molecule optionally comprise another peptide, e.g. a peptide facilitating purification, in particular a peptide being an epitope against which an antibody is available, such as the FLAG peptide. Purification, e.g. by means of affinity chromatography, of a fusion protein comprising such a peptide is advantageous e.g. in that it may be faster, more specific and/or gentler. The peptide may be placed at the N-terminus of the fusion protein, in between the recombinant antibody and the effector molecule, or at the C-terminus of the fusion protein. Preferably, it is located at the N-terminus or at the C-terminus, in particular at the N-terminus. Preferably, these constructs also contain a cleavage site, so that the fusion protein can be liberated therefrom, either by enzymatic cleavage, e.g. by enterokinase or by Factor Xa, or by the chemical methods known in the art. Furthermore these constructs may comprise a peptide spacer consisting of one or more, e.g. 1 to 10, in particular about 2 amino acids, said spacer facilitating the linkage of the above-mentioned peptide and/or the cleavage site to the recombinant antibody. The cleavage site is placed in such a way that the fusion protein comprising the recombinant antibody and the effector molecule can be easily liberated, if desired, preferably in vitro. For example, in the protein construct comprising the fusion protein designated Fv(FRP5)-ETA (cf. SEQ. ID NO: 10), the FLAG peptide and an enterokinase cleavage site are linked to a spacer and placed in front of the Fv heavy chain/light chain variable domain and exotoxin A fusion protein. If desired, the FLAG peptide can be cleaved off by enterokinase, preferably after affinity purification of the protein, yielding a fusion protein comprising the single-chain antibody Fv(FRP5) and exotoxin

Most preferred is a single-chain recombinant antibody wherein the heavy chain variable domain and the light chain variable domain are derived from a mouse monoclonal antibody directed to the extracellular domain of the growth factor receptor c-erbB-2, e.g. derived from the mouse monoclonal antibodies FRP5, FSP16, FWP51 or FSP77, particularly from the mouse monoclonal antibodies FRP5 or FWP51. Likewise preferred is a single-chain recombinant antibody wherein the spacer group linking the light chain and the heavy chain variable domains is a polypeptide comprising about 15 amino acids selected from glycine and serine, in particular wherein the spacer group is the 15 amino acid polypeptide consisting of three repetitive subunits of Gly-Gly-Gly-Ser.

Especially preferred is a single-chain antibody comprising the heavy chain variable domain of a mouse monoclonal antibody selected from the group consisting of FRP5, FSP16, FWP51 and FSP77, the 15 amino acid spacer group consisting of three repetitive subunits of Gly-Gly-Gly-Gly-Ser, the light chain variable domain of a mouse monoclonal antibody selected from the group consisting of FRP5, FSP16, FWP51 and FSP77 and an enzyme, for example a phosphatase such as the alkaline phosphatase phoA, or an exotoxin such as Pseudomonas exotoxin, or a variant thereof.

Particularly preferred is the particular single-chain recombinant antibody designated Fv(FRP5)-phoA comprising a polypeptide of the amino acid sequence 2 to 690 of SEQ ID NO: 5.

Likewise preferred is a single-chain recombinant antibody comprising a peptide facilitating purification, a cleavage site and a particular single-chain recombinant antibody selected from the group consisting of Fv(FRPS)-ETA and Fv (FWP51)-ETA, in particular a single-chain recombinant antibody comprising a polypeptide selected from the group consisting of a polypeptide of the amino acid sequence -10 to 606 of SEQ. ID NO:10 and of a polypeptide of the amino acid sequence -10 to 606 of SEQ. ID NO:11, said protein being subjected to in vitro cleavage by enterokinase, if desired.

Particularly preferred is a single-chain recombinant antibody comprising a protein selected from the group consisting of a polypeptide of the amino acid sequence 2 to 606 of SEQ ID NO:10 and a polypeptide of the amino acid sequence 2 to 606 of SEQ ID NO:11.

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The invention further concerns the mouse monoclonal antibodies directed to the extracellular domain of the growth factor receptor c-erbB-2 and designated FRPS, FSP16, FSP77, and FWP51, which are secreted by the hybridoma cell lines FRPS, FSP16, FSP77, and FWP51, respectively. Most preferred are the mouse monoclonal antibodies designated FRPS and FWP51.

The invention further concerns a method of manufacture of the recombinant antibodies and of the mouse monoclonal antibodies of the invention. The antibodies are prepared by processes that are known <u>per se</u>, characterized in that host cells or hybridoma cells as defined further below producing such antibodies are multiplied <u>in vitro</u> or <u>in vivo</u> and, when required, the obtained antibodies are isolated. For example, the recombinant antibodies of the invention can be prepared by recombinant DNA techniques comprising culturing a transformed host under conditions which allow expression thereof and isolating said antibody.

More specifically, the present invention also relates to a process for the production of a protein of the invention selected from the group consisting of a heavy chain murine variable domain, a light chain murine variable domain, a heavy chain murine variable domain and a light chain murine variable domain, a single-chain recombinant antibody, a fusion protein, and a fusion protein optionally comprising a peptide facilitating purification, a cleavage site and a peptide spacer comprising culturing a host, e.g. E. coli, which has been transformed with a hybrid vector comprising an expression cassette comprising a promoter and a DNA coding for said protein which DNA is controlled by said promoter, and isolating said protein.

In particular, the present invention relates to a process for the production of a protein of the invention selected from the group consisting of a heavy chain murine variable domain, a light chain murine variable domain, a heavy chain murine variable domain and a light chain murine variable domain, a single-chain recombinant antibody, and a fusion protein optionally comprising a peptide facilitating purification, a cleavage site and a peptide spacer comprising culturing a host, e.g. E. coli, which has been transformed with a hybrid vector comprising an expression cassette comprising a promoter operably linked to a first DNA sequence encoding a signal peptide linked in the proper reading frame to a second DNA sequence encoding said protein, and isolating said protein.

Multiplication of hybridoma cells or mammalian host cells in vitro is carried out in suitable culture media, which are the customary standard culture media, for example Dulbecco's Modified Eagle Medium (DMEM) or RPMI 1640 medium, optionally replenished by a mammalian serum, e.g. fetal calf serum, or trace elements and growth sustaining supplements, e.g feeder cells such as normal mouse peritoneal exudate cells, spleen cells, bone marrow macrophages, 2-aminoethanol, insulin, transferrin, low density lipoprotein, oleic acid, or the like. Multiplication of host cells which are bacterial cells or yeast cells is likewise carried out in suitable culture media known in the art, for example for bacteria in medium LB, NZCYM, NZYM, NZYM, Terrific Broth, SOB, SOC, 2 x YT, or M9 Minimal Medium, and for yeast in medium YPD, YEPD, Minimal Medium, or Complete Minimal Dropout Medium.

<u>In vitro</u> production provides relatively pure antibody preparations and allows scale-up to give large amounts of the desired antibodies. Techniques for bacterial cell, yeast or mammalian cell cultivation are known in the art and include homogeneous suspension culture, e.g. in an airlift reactor or in a continuous stirrer reactor, or immobilized or entrapped cell culture, e.g. in hollow fibres, microcapsules, on agarose microbeads or ceramic cartridges.

Large quantities of the desired antibodies can also be obtained by multiplying mammalian cells in vivo. For this purpose, hybridoma cells producing the desired antibodies are injected into histocompatible mammals to cause growth of antibody-producing tumors. Optionally, the animals are primed with a hydrocarbon, especially mineral oils such as pristane (tetramethyl-pentadecane), prior to the injection. After one to three weeks, the antibodies are isolated from the body fluids of those mammals. For example, hybridoma cells obtained by fusion of suitable myeloma cells with antibody-producing spleen cells from Balb/c mice, or transfected cells derived from hybridoma cell line Sp2/0 that produce the desired antibodies are injected intraperitoneally into Balb/c mice optionally pre-treated with pristane, and, after one to two weeks, ascitic fluid is taken from the animals.

The cell culture supernatants are screened for the desired antibodies, preferentially by immunofluorescent staining of cells expressing c-erbB-2, by immunoblotting, by an enzyme immunoassay, e.g. a sandwich assay or a dot-assay, or a radioimmunoassay.

For isolation of the antibodies, the immunoglobulins in the culture supernatants or in the ascitic fluid may be concentrated, e.g. by precipitation with ammonium sulphate, dialysis against hygroscopic material such as polyethylene glycol, filtration through selective membranes, or the like. If necessary and/or desired, the antibodies are purified by the customary chromatography methods, for example gel filtration, ion-exchange chromatography, chromatography over DEAE-cellulose and/or (immuno-)affinity chromatography, e.g. affinity chromatography with c-erbB-2 protein or with Protein-A.

The invention further concerns hybridoma cells secreting the monoclonal antibodies of the invention, in particular the hybridoma cell lines FRP5, FSP16, FSP77, and FWP51 deposited under the Budapest Treaty on November 21,

1990 at the European Collection of Animal Cell Cultures (ECACC) in Porton Down, Salibury, UK, under the accession numbers 90112115, 90112116, 90112117, and 90112118, respectively. Most preferred is the hybridoma cell line designated FRP5, ECACC number 90112115 or the hybridoma cell line designated FWP51, ECACC number 90112118. The preferred hybridoma cells of the invention are genetically stable, secrete monoclonal antibodies of the invention of the desired specificity and can be activated from deep-frozen cultures by thawing and recloning.

The invention also concerns a process for the preparation of a hybridoma cell line secreting monoclonal antibodies directed to the extracellular domain of the growth factor receptor c-erbB-2, characterized in that a suitable mammal, for example a Balb/c mouse, is immunized with purified c-erbB-2 protein, an antigenic carrier containing purified c-erbB-2 or with cells bearing growth factor receptor c-erbB-2, antibody-producing cells of the immunized mammal are fused with cells of a suitable myeloma cell line, the hybrid cells obtained in the fusion are cloned, and cell clones secreting the desired antibodies are selected. For example spleen cells of Balb/c mice immunized with cells bearing c-erbB-2 are fused with cells of the myeloma cell line PAI or the myeloma cell line Sp2/0-Ag 14, the obtained hybrid cells are screened for secretion of the desired antibodies, and positive hybridoma cells are cloned.

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Preferred is a process for the preparation of a hybridoma cell line, characterized in that Balb/c mice are immunized by injecting subcutaneously and/or intraperitoneally between 10⁷ and 10⁸ cells of the human breast tumor cell line SKBR3 containing a suitable adjuvant several times, e.g. four to six times, over several months, e.g. between two and four months, and spleen cells from the immunized mice are taken two to four days after the last injection and fused with cells of the myeloma cell line PAI in the presence of a fusion promoter, preferably polyethylene glycol. Preferably the myeloma cells are fused with a three- to twentyfold excess of spleen cells from the immunized mice in a solution containing about 30 % to about 50 % polyethylene glycol of a molecular weight around 4000. After the fusion the cells are expanded in suitable culture media as described hereinbefore, supplemented with a selection medium, for example HAT medium, at regular intervals in order to prevent normal myeloma cells from overgrowing the desired hybridoma cells.

The invention also concerns recombinant DNAs comprising an insert coding for a heavy chain murine variable domain and/or for a light chain murine variable domain of antibodies directed to the extracellular domain of the growth factor receptor c-erbB-2 as described hereinbefore. By definition such DNAs comprise coding single stranded DNAs, double stranded DNAs consisting of said coding DNAs and of complementary DNAs thereto, or these complementary (single stranded) DNAs themselves.

Furthermore, DNA encoding a heavy chain murine variable domain and/or for a light chain murine variable domain of antibodies directed to the extracellular domain of the growth factor receptor c-erbB-2 can be enzymatically or chemically synthezised DNA having the authentic DNA sequence coding for a heavy chain murine variable domain and/or for the light chain murine variable domain, or a mutant therof. A mutant of the authentic DNA is a DNA encoding a heavy chain murine variable domain and/or a light chain murine variable domain of the above-mentioned antibodies in which one or more amino acids are deleted or exchanged with one or more other amino acids. Preferably said modification(s) are outside the CDRs of the heavy chain murine variable domain and/or of the light chain murine variable domain of the antibody. Such a mutant DNA is also intended to be a silent mutant wherein one or more nucleotides are replaced by other nucleotides with the new codons coding for the same amino acid(s). Such a mutant sequence is also a degenerated sequence. Degenerated sequences are degenerated within the meaning of the genetic code in that an unlimited number of nucleotides are replaced by other nucleotides without resulting in a change of the amino acid sequence originally encoded. Such degenerated sequences may be useful due to their different restriction sites and/or frequency of particular codons which are preferred by the specific host, particularly E. coli, to obtain an optimal expression of the heavy chain murine variable domain and/or a light chain murine variable domain.

The term mutant is intended to include a DNA mutant obtained by in vitro mutagenesis of the authentic DNA according to methods known in the art.

The invention relates to a recombinant DNA comprising an insert coding for a heavy chain murine variable domain of a monoclonal antibody selected from the group consisting of the antibodies FRP5, FSP16, FSP77 and FWP51, or coding for an amino acid sequence homologous to said heavy chain variable domain.

In particular, the invention concerns a recombinant DNA comprising an insert coding for a heavy chain murine variable domain, which originates from genomic DNA or mRNA of the hybridoma cell lines FRP5, FSP16, FSP77 or FWP51, or which is homologous to genomic DNA of said cell lines and codes for an amino acid sequence homologous to the heavy chain variable domain of monoclonal antibodies FRP5, FSP16, FSP77 or FWP51. Especially preferred is a recombinant DNA comprising an insert coding for a heavy chain murine variable domain, which originates from genomic DNA or mRNA of the hybridoma cell line FRP5, or which is homologous to genomic DNA of said cell line and codes for an amino acid sequence homologous to the heavy chain variable domain of monoclonal antibody FRP5; or a recombinant DNA or mRNA of the hybridoma cell line FWP51, or which is homologous to genomic DNA of said cell line and codes for an amino acid sequence homologous to the heavy chain variable domain of monoclonal antibody FWP51.

Preferred is a recombinant DNA comprising an insert coding for the polypeptide of formula I, wherein FR₁, FR₂,

FR₃, FR₄, CDR_{1H}, CDR_{2H}, and CDR_{3H} have the meanings as mentioned hereinbefore, optionally further containing introns. Especially preferred is a recombinant DNA coding for the polypeptide of formula I comprising inserts coding for murine or human framework regions FR₁, FR₂, FR₃ and FR₄, and inserts coding for complementarity determining regions of the DNA sequence 99 to 113 (CDR_{1H}), the DNA sequence 156 to 206 (CDR_{2H}), and the DNA sequence 303 to 332 (CDR_{3H}) of SEQ ID NO:4 or coding for complementarity determining regions of the DNA sequence 99 to 113 (CDR_{1H}), the DNA sequence 156 to 206 (CDR_{2H}), and the DNA sequence 303 to 335 (CDR_{3H}) of SEQ ID NO:8. Most preferred is a DNA comprising an insert of the DNA sequence 9 to 365 of SEQ ID NO:4, wherein optionally one or more, e.g. 1 to 10, nucleotides are replaced by other nucleotides, in particular a DNA comprising an insert of the DNA sequence 9 to 365 of SEQ ID NO:8, wherein optionally one or more, e.g. 1 to 10, nucleotides are replaced by other nucleotides are replaced by other nucleotides, in particular a DNA comprising an insert of the DNA sequence 9 to 368 of SEQ ID NO:8, wherein optionally one or more, e.g. 1 to 10, nucleotides are replaced by other nucleotides, in particular a DNA comprising an insert of the DNA sequence 9 to 368 of SEQ ID NO:8.

In a DNA wherein nucleotides of the sequence given in SEQ ID NO:4, or in a DNA wherein nucleotides of the sequence given in SEQ ID NO:8, are replaced by other nucleotides, such replacement is preferred when it does not alter the amino acid sequence of the complementarity determining regions (CDRs) coded for. This means that such replacement of nucleotides may occur in the inserts coding for the framework regions (FRs) or in a position where it does not alter the amino acid coded for due to the degeneracy of the triplet codons.

Likewise the invention relates to a recombinant DNA comprising an insert coding for a light chain murine variable domain of a monoclonal antibody selected from the group consisting of the antibodies FRP5, FSP16, FSP77 and FWP51, or coding for an amino acid sequence homologous to said light chain variable domain.

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More specifically, the invention concerns a recombinant DNA comprising an insert coding for a light chain murine variable domain, which originates from genomic DNA or mRNA of the hybridoma cell lines FRP5, FSP16, FSP77 or FWP51, or which is homologous to genomic DNA of said cell lines and codes for an amino acid sequence homologous to the light chain variable domain of monoclonal antibodies FRP5, FSP16, FSP77 or FWP51. Particularly preferred is a recombinant DNA comprising an insert coding for a light chain murine variable domain, which originates from genomic DNA or mRNA of the hybridoma cell line FRP5, or which is homologous to genomic DNA of said cell line and codes for an amino acid sequence homologous to the light chain variable domain of monoclonal antibody FRP5, or a recombinant DNA comprising an insert coding for a light chain murine variable domain, which originates from genomic DNA or mRNA of the hybridoma cell line FwP51, or which is homologous to genomic DNA of said cell line and codes for an amino acid sequence homologous to the light chain variable domain of monoclonal antibody FWP51.

Preferred is a recombinant DNA comprising an insert coding for the polypeptide of formula II, wherein FR₅, FR₆, FR₇, FR₈, CDR_{1L}, CDR_{2L}, and CDR_{3L} have the meanings as mentioned hereinbefore, optionally further containing introns. Especially preferred is a recombinant DNA coding for the polypeptide of formula II comprising inserts coding for murine or human framework regions FR₅, FR₆, FR₇ and FR₈, and inserts coding for complementarity determining regions of the DNA sequence 480 to 512 (CDR_{1L}), the DNA sequence 558 to 578 (CDR_{2L}), and the DNA sequence 675 to 701 (CDR_{3L}) of SEQ ID NO:4, or coding for complementarity determining regions of the DNA sequence 483 to 515 (CDR_{1L}), the DNA sequence 561 to 581 (CDR_{2L}), and the DNA sequence 678 to 701 (CDR_{3L}) of SEQ ID NO:8.

Most preferred is a DNA comprising an insert of the DNA sequence 411 to 728 of SEQ ID NO:4, wherein optionally one or more, e.g. 1 to 10, nucleotides are replaced by other nucleotides, in particular a DNA comprising an insert of the DNA sequence 411 to 728 of SEQ ID NO:4. Likewise preferred is a DNA comprising an insert of the DNA sequence 414 to 728 of SEQ ID NO:8, wherein optionally one or more, e.g. 1 to 10, nucleotides are replaced by other nucleotides, in particular a DNA comprising an insert of the DNA sequence 414 to 728 of SEQ ID NO:8. In a DNA wherein nucleotides of the sequence given in SEQ ID NO:4, or in a DNA wherein nucleotides of the sequence given in SEQ ID NO:8, are replaced by other nucleotides, such replacement is preferred when it does not after the amino acid sequence of the complementarity determining regions (CDRs) coded for, as is described above for DNA coding for the heavy chain variable domain

For the assembly of complete tetrameric immunoglobulin molecules and the expression of chimeric antibodies, the recombinant DNA inserts coding for heavy and light chain variable domains are fused with the corresponding DNAs coding for heavy and light chain constant domains, then transferred into appropriate host cells, for example after incorporation into hybrid vectors.

The invention therefore also concerns recombinant DNAs comprising an insert coding for a heavy chain murine variable domain of an antibody directed to the extracellular domain of c-erbB-2 fused to a human constant domain γ , for example γ 1, γ 2, γ 3 or γ 4, preferably γ 1 or γ 4. Likewise the invention concerns recombinant DNAs comprising an insert coding for a light chain murine variable domain of an antibody directed to the extracellular domain of c-erbB-2 fused to a human constant domain κ or λ , preferably κ .

The invention especially concerns recombinant DNAs coding for a single-chain recombinant antibody as defined hereinbefore, e.g. recombinant DNA wherein the heavy chain variable domain and the light chain variable domain are linked by way of a DNA insert coding for a spacer group, in particular a recombinant DNA coding for a protein of the formula III, wherein FR₁,FR₂,FR₃,FR₄,FR₆,FR₇,FR₈,FR₉, SP, CDR_{1H}, CDR_{2H}, CDR_{3H}, CDR_{3L}, CDR_{2L} and CDR_{3L} have

the meanings given above, optionally comprising further DNA coding for an effector molecule and/or signal sequences facilitating the processing of the antibody in the host cell. In particular the invention concerns a DNA comprising an insert of the DNA sequence 9-728 of SEQ ID NO:4, wherein optionally one or more, e.g. 1 to 10, nucleotides are replaced by other nucleotides, especially a DNA comprising an insert of the DNA sequence 9 to 728 of SEQ ID NO:4. Furthermor the invention relates to a DNA comprising an insert of the DNA sequence 9-728 of SEQ ID NO:8 wherein optionally one or more, e.g. 1 to 10, nucleotides are replaced by other nucleotides, especially a DNA comprising an insert of the DNA sequence 9 to 728 of SEQ ID NO:8.

In another embodiment the invention pertains to recombinant DNAs coding for a recombinant DNA wherein the heavy chain variable domain and the light chain variable domain are linked by way of a DNA insert coding for a spacer group, optionally comprising a signal sequence facilitating the processing of the antibody in the host cell and/or a DNA coding for a peptide facilitating the purification of the antibody and/or a DNA coding for a cleavage site and/or a DNA coding for a peptide spacer and/or a DNA coding for an effector molecule.

The DNA coding for an effector molecule is intended to be a DNA coding for the above-mentioned effector molecules, particularly a DNA coding for alkaline phosphatase or Pseudomonas exotoxin A. The DNA encoding such an effector molecule has the sequence of a naturally occurring enzyme or toxin encoding DNA, or a mutant therof, and can be prepared by methods well known in the art. A mutant of the naturally occurring DNA encoding e.g. alkaline phosphatase or Pseudomonas exotoxin A, or a variant thereof can be obtained e.g. analogously to the methods described above.

Most preferred is a DNA comprising an insert of the DNA sequence 23 to 814 of SEQ ID NO:5, of the DNA sequence 86 to 2155 of SEQ ID NO:5 or of the DNA sequence 23 to 2155 of SEQ ID NO:5, wherein optionally one or more, e. g. 1 to 10, nucleotides are replaced by other nucleotides, in particular a DNA comprising an insert of the DNA sequence 23 to 2155 of SEQ ID NO:5.

Equally preferred is a DNA comprising an insert of the DNA sequence 1 to 1911 of SEQ ID NO: 10, of the DNA sequence 64 to 1911 of SEQ ID NO: 10, or of the DNA sequence 97 to 1911 of SEQ ID NO: 10, wherein optionally one or more, e.g. 1 to 10, nucleotides are replaced by other nucleotides, in particular a DNA comprising an insert of the DNA sequence 1 to 1911 of SEQ IDs NO: 10; or a DNA comprising an insert of the DNA sequence 1 to 1911 of SEQ ID NO: 11, of the DNA sequence 96 to 1911 of SEQ ID NO: 11, or of the DNA sequence 97 to 1911 of SEQ ID NO: 11, wherein optionally one or more, e.g. 1 to 10, nucleotides are replaced by other nucleotides, in particular a DNA comprising an insert of the DNA sequence 1 to 1911 of SEQ ID NO: 11.

Furthermore the invention concerns a recombinant DNA which is a hybrid vector comprising an insert coding for the variable domain of a murine heavy chain as described hereinbefore and/or an insert coding for the variable domain of a murine light chain as described hereinbefore, an origin of replication or an autonomously replicating sequence, one or more dominant marker sequences and, optionally, expression control sequences, signal sequences and additional restriction sites.

In a first embodiment the hybrid vector according to the invention comprises an expression cassette comprising a promoter and a DNA coding for a protein of the invention selected from the group consisting of a heavy chain murine variable domain, a light chain murine variable domain, a light chain murine variable domain, a single-chain recombinant antibody, a fusion protein, and a fusion protein optionally comprising a peptide facilitating purification, a cleavage site and a peptide spacer, which DNA is controlled by said promoter, and isolating said protein.

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In a second embodiment, the hybrid vector according to the invention comprises an expression cassette comprising a promoter operably linked to a first DNA sequence encoding a signal peptide linked in the proper reading frame to a second DNA sequence encoding a protein of the invention selected from the group consisting of a heavy chain murine variable domain, a light chain murine variable domain, a light chain murine variable domain, a single-chain recombinant antibody, and a fusion protein optionally comprising a peptide facilitating purification, a cleavage site and a peptide spacer.

Vectors typically perform two functions in collaboration with compatible host cells. One function is to facilitate the cloning of the nucleic acid that encodes the immunoglobulin variable domains, i.e. to produce usable quantities of the nucleic acid (cloning vectors). The other function is to provide for replication and expression of the recombinant gene constructs in a suitable host, either by maintenance as an extrachromosomal element or by integration into the host chromosome (expression vectors). A cloning vector comprises the recombinant gene constructs as described above, an origin of replication or an autonomously replicating sequence, dominant marker sequences and, optionally, signal sequences and additional restriction sites. An expression vector additionally comprises expression control sequences essential for the transcription and translation of the recombinant genes.

An origin of replication or an autonomously replicating sequence is provided either by construction of the vector to include an exogeneous origin such as derived from Simian virus 40 (SV 40) or another viral source, or by the host cell chromosomal mechanisms.

The markers allow for selection of host cells which contain the vector. Selection markers include genes which

confer resistance to heavy metals such as copper or to antibiotics such as geneticin (G-418) or hygromycin, or genes which complement a genetic lesion of the host cell such as the absence of thymidin kinase, hypoxanthine phosphoryl transferase, dihydrololate reductase or the like.

Signal sequences may be, for example, presequences or secretory leaders directing the secretion of the recombinant antibody, splice signals, or the like. Examples for signal sequences directing the secretion of the recombinant antibody are sequences derived from the ompA gene, the pelB (pectate lyase) gene or the phoA gene.

As expression control sequences, the vector DNA comprises a promoter, sequences necessary for the initiation and termination of transcription and for stabilizing the mRNA and, optionally, enhancers and further regulatory sequences.

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A wide variety of promoting sequences may be employed, depending on the nature of the host cell. Promoters that are strong and at the same time well regulated are the most useful. Sequences for the initiation of translation are for example Shine-Dalgarno sequences. Sequences necessary for the initiation and termination of transcription and for stabilizing the mRNA are commonly available from the noncoding 5'-regions and 3'-regions, respectively, of viral or eukaryotic cDNAs, e.g. from the expression host. Enhancers are transcription-stimulating DNA sequences of viral origin, e.g. derived from Simian virus, polyoma virus, bovine papilloma virus or Moloney sarcoma virus, or of genomic, especially murine, origin.

The various DNA segments of the vector DNA are operationally linked, i.e. they are contiguous and placed into a functional relationship with each other.

Examples of vectors which are suitable for replication and expression in an \underline{E} . \underline{coli} strain are bacteriophages, for example derivatives of λ bacteriophages, or plasmids, such as, in particular, the plasmid ColE1 and its derivatives, for example pMB9, pSF2124, pBR317 or pBR322 and plasmids derived from pBR322, such as pUC9, pUCK0, pHRi148 and pLc24. Suitable vectors contain a complete replicon, a marker gene, recognition sequences for restriction endonucleases, so that the foreign DNA and, if appropriate, the expression control sequence can be inserted at these sites, and optionally signal sequences and enhancers.

Microbial promoters are, for example, the strong leftward promoter P_L of bacteriophage λ which is controlled by a temperature sensitive repressor. Also suitable are \underline{E} . \underline{coli} promoters such as the lac (lactose) promoter regulated by the lac repressor and induced by isopropyl- β -D-thiogalactoside, the trp (tryptophan) promoter regulated by the trp repressor and induced e.g. by tryptophan starvation, and the tac (hybrid trp-lac promoter) regulated by the lac repressor.

Vectors which are suitable for replication and expression in yeast contain a yeast replication start and a selective genetic marker for yeast One group of such vectors includes so-called ars sequences (autonomous replication sequences) as origin of replication. These vectors are retained extrachromosomally within the yeast cell after the transformation and are replicated autonomously. Furthermore, vectors which contain all or part of the 2µ (2 mikron) plasmid DNA from Saccharomyces cerevisiae can be used. Such vectors will get integrated by recombination into 2µ plasmids already existing within the cell, or replicate autonomously. 2µ sequences are particularly suitable when high transformation frequency and high copy numbers are to be achieved.

Expression control sequences which are suitable for expression in yeast are, for example, those of highly expressed yeast genes. Thus, the promoters for the <u>TRP1</u> gene, the <u>ADHI</u> or <u>ADHII</u> gene, acid phosphatase (<u>PHO3</u> or <u>PHO5</u>) gene, isocytochrome gene or a promoter involved with the glycolytic pathway, such as the promoter of the enolase, glyceraldehyde-3-phosphate kinase (<u>PGK</u>), hexokinase, pyruvate decarboxylase, phosphofructokinase, glucose-6-phosphate isomerase, 3-phosphoglycerate mutase, pyruvate kinase, triosephosphate isomerase, phosphoglycese isomerase and glucokinase genes, can be used.

Vectors suitable for replication and expression in mammalian cells are preferably provided with promoting sequences derived from DNA of viral origin, e.g. from Simian virus 40 (SV40), Rous sarcoma virus (RSV), adenovirus 2, bovine papilloma virus (BPV), papovavirus BK mutant (BKV), or mouse or human cytomegalovirus (CMV). Alternatively, the vectors may comprise promoters from mammalian expression products, such as actin, collagen, myosin etc., or the native promoter and control sequences which are normally associated with the desired gene sequence, i.e. the immunoglobulin H-chain or L-chain promoter.

Preferred vectors are suitable for both procaryotic and eucaryotic hosts and are based on viral replication systems. Particularly preferred are vectors comprising Simian virus promoters, e.g. pSVgpt or pSVneo, further comprising an enhancer, e.g. an enhancer normally associated with the immunoglobulin gene sequences, in particular the mouse Ig H- or L-chain enhancer.

The recombinant DNA coding for a recombinant antibody of the invention can be prepared, for example, by culturing a transformed host cell and optionally isolating the prepared DNA.

In particular, such DNA can be prepared by a method comprising

a) preparing murine DNA coding for the variable heavy and/or light chain domains of the antibody with the desired specificity, e.g. by isolating the DNA from the genome of a suitable hybridoma cell line and selecting the desired DNA using DNA probes, or by isolating mRNA from a suitable hybridoma cell line and preparing cDNA coding for the variable heavy and/or light chain domains of the antibody with the desired specificity using oligonucleotide primers.

- b) preparing DNA coding for the desired signal sequence and/or preparing DNA coding for an effector molecule,
 e.g. by isolating the desired DNA(s) from a suitable source,
 e.g. from a genomic library or a cDNA library using DNA probes.
- c) synthesizing DNA coding for the desired spacer group by chemical methods,
- d) constructing recombinant genes encoding the recombinant antibodies by incorporating the DNA of step a) and, optionally, b) and/or c) into appropriate hybrid vectors,
- e) transferring the obtained hybrid vectors into a recipient host cell or retrieving the DNA coding for the recombinant genes and transferring the unlinked DNA into a recipient host cell,
- f) selecting and culturing the transformed host cell, and
- g) optionally isolating the desired DNA.

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The DNA according to step a) of the process described above can be obtained by isolation of genomic DNA or by preparation of cDNA from isolated mRNA. Genomic DNA from hybridoma cells is isolated by methods known in the art which include steps for disruption of the cells, e.g. by lysis in presence of detergents like Triton™, extracting the DNA, e.g. by treatment with phenol and CHCl₃/isoamyl alcohol, and precipitation of DNA. The DNA is fragmented, conveniently by one or more restriction endonucleases, the resulting fragments are replicated on a suitable carrier, e. g. nitrocellulose membranes, and screened with a DNA probe for the presence of the DNA sequences coding for the polypeptide sequence of interest, in particular for the presence of the rearranged H- and L-chain Ig gene loci. By this procedure DNA fragments are found that contain inserts with heavy chain V, D and J regions and light chain V and J regions, respectively, together with a leader sequence and introns, if any. cDNA from hybridoma cells is likewise prepared by methods known in the art, e.g. by extracting total cellular RNA, isolating mRNA by a suitable chromatographic method, e.g. chromatography on oligo(dT)-cellulose, synthesizing cDNA with a mixture of deoxynucleotide triphosphates and reverse transcriptase in the presence of oligonucleotide primers complementary to suitable regions in the murine immunoglobulin heavy and light chain constant domain genes, and isolating the cDNA. As a tool simplifying DNA isolation, the desired genomic DNA or cDNA may be amplified using polymerase chain reaction (PCR) technology. PCR involves repeated rounds of extension from two primers specific for DNA regions at each end of the gene. Preferably, cDNA transcripts of total mRNA from the suitable hybridoma cell line is treated in a heating/cooling cycle with Tag DNA polymerase in the presence of primers tailored to hybridize to Ig H- and L-chain variable domains, respectively.

Genomic DNA or cDNA according to step b) of the process described above is isolated from suitable bacterial or mammalian cells according to methods known in the art. Preferably, the methods as described under a) are used, substituting the corresponding source cells for the murine hybridoma cells and using DNA probes designed to hybridize with the desired signal sequences or the genes coding for the desired effector molecules. In bacteria wherein separation of mRNA from total RNA is not possible with olig(dT)-cellulose, cDNA is prepared from total RNA using corresponding oligonucleotide primers. The DNA isolation is simplified considerably by the PCR technology.

DNA according to step c) is prepared by conventional chemical and enzymatic methods, e.g. by chemical synthesis of oligonucleotides of between thirty and sixty bases with overlapping complementary sequences, hybridization of such oligonucleotides, and enzymatic ligation, optionally after filling-in of missing bases with suitable enzymes in the presence of the corresponding deoxynucleotide triphosphates.

The DNA probe for the mouse variable chain domains may be a synthetic DNA, a cDNA derived from mRNA coding for the desired immunoglobulin or a genomic DNA or DNA fragment of known nucleotide sequence. As probes for the detection and/or amplification of the rearranged lg gene loci of the variable domains of L-/H-chains, DNA fragments of known nucleotide sequences of adjacent conserved variable or constant domains are selected which constitute the lg loci of the L-/H-chain in the mammal from which the DNA is derived, e.g. Balb/c mice. The DNA probe is synthesized by chemical methods or isolated from suitable tissue of an appropriate mammal, e.g. Balb/c mouse liver, and purified by standard methods. If required, the probe DNA is labelled, e.g. radioactively labelled by the well-known nick-translation technique, then hybridized with the DNA library in buffer and salt solutions containing adjuncts, e.g. calcium chelators, viscosity regulating compounds, proteins, non-specific DNA and the like, at temperatures favoring selective hybridization.

Once a fragment has been identified which contains the desired DNA sequence, this fragment may be further manipulated to remove nonessential DNA, modified at one or both termini, and treated to remove all or a portion of intervening sequences, or the like.

The joining of the various DNA fragments in order to produce recombinant genes encoding the recombinant antibodies is performed in accordance with conventional techniques, for example, by blunt- or staggered-end ligation, restriction enzyme digestion to provide for appropriate cohesive termini, filling-in cohesive ends as appropriate, alkaline phosphatase treatment to avoid undesirable joining, and ligation with appropriate ligases.

The transfer of the recombinant DNAs, e.g. the transfer of hybrid vectors, and the selection of transformed cells

is described below.

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Moreover, the invention relates to host cells transformed with the recombinant DNAs described above, namely host cells which are transformed with a DNA encoding the heavy chain and/or a DNA encoding the light chain of the desired recombinant antibody, in particular host cells transformed with a DNA encoding the preferred single-chain recombinant antibody.

More specifically, the invention concerns a host cell which has been transformed with a hybrid vector comprising an expression cassette comprising a promoter and a DNA coding for a protein of the invention selected from the group consisting of a heavy chain murine variable domain, a light chain murine variable domain, a heavy chain murine variable domain and a light chain murine variable domain, a single-chain recombinant antibody, a fusion protein, and a fusion protein further comprising a peptide facilitating purification, a cleavage site and a peptide spacer which DNA is controlled by said promoter.

Furthermore, the invention pertains to a host cell which has been transformed with a hybrid vector comprising an expression cassette comprising a promoter operably linked to a first DNA sequence encoding a signal peptide linked in the proper reading frame to a second DNA sequence encoding a protein of the invention selected from the group consisting of a heavy chain murine variable domain, a light chain murine variable domain, a heavy chain murine variable domain and a light chain murine variable domain, a single-chain recombinant antibody, a fusion protein, and a fusion protein further comprising a peptide facilitating purification, a cleavage site and a peptide spacer.

In particular, the present invention relates to a process for the production of a protein of the invention selected from the group consisting of a heavy chain murine variable domain, a light chain murine variable domain, a heavy chain murine variable domain and a light chain murine variable domain, a single-chain recombinant antibody, a fusion protein, and a fusion protein further comprising a peptide facilitating purification, a cleavage site and a peptide spacer comprising culturing a host, e.g. E. coli, which has been transformed with a hybrid vector comprising an expression cassette comprising a promoter operably linked to a first DNA sequence encoding a signal peptide linked in the proper reading frame to a second DNA sequence encoding said protein, and isolating said protein.

The host cells of the present invention have to be capable of culture in vitro. Suitable host cells are of procaryotic or of eucaryotic origin and are, for example, bacterial cells, e.g. E. coli, yeasts, e.g. Saccharomyces cerevisiae, or mammalian cells. For the preparation of functional chimeric human/mouse antibodies the host cells have to be of higher eucaryotic origin to provide a suitable environment for the production of active antibodies, since the biosynthesis of functional tetrameric antibody molecules requires correct nascent polypeptide chain folding, glycosylation, and assembly.

Examples of suitable hosts are microorganisms which are devoid of or poor in restriction enzymes or modification enzymes, such as bacteria, in particular strains of Escherichia coli, for example E. coli X1776, E. coli Y1090, E. coli HB 101, E. coli W3110, E. coli HB101/LM1035, E. coli JA 221, E. coli DH5α, E. coli K12, or E. coli CC118 strain, Bacillus subtilis, Bacillus stearothermophilus, Pseudomonas, Haemophilus, Streotococcus and others, and yeasts, for example Saccharomyces cerevisiae such as S. cerevisiae GRF 18. Further suitable host cells are cells of higher organisms, in particular established continuous human or animal cell lines, e.g. human embryonic lung fibroblasts L132, human malignant melanoma Bowes cells, HeLa cells, SV40 virus transformed kidney cells of African green monkey COS-7 or Chinese hamster ovary (CHO) cells, or cells of lymphoid origin, such as lymphoma, myeloma, hybridoma, trioma or quadroma cells, for example PAI, Sp2/0 or X63-Ag8.653 cells.

The above mentioned strains of \underline{E} . \underline{coli} , in particular \underline{E} . \underline{coli} CC118, are preferred as hosts.

The invention also concerns processes for the preparation of transformed host cells wherein suitable recipient host cells as described hereinbefore are transformed with a hybrid vector according to the invention, and the transformed cells are selected.

Transformation of microorganisms is carried out as described in the literature, for example for <u>S. cerevisiae</u> (A. Hinnen et al., Proc. Natl. Acad. Sci. USA 75: 1929, 1978), for <u>B. subtilis</u> (Anagnostopoulos et al., J. Bacteriol. 81: 741, 1961), and for <u>E. coli</u> (M. Mandel et al., J. Mol. Biol. 53: 159, 1970).

Accordingly, the transformation procedure of <u>E. coli</u> cells includes, for example, Ca²⁺ pretreatment of the cells so as to allow DNA uptake, and incubation with the hybrid vector. The subsequent selection of the transformed cells can be achieved, for example, by transferring the cells to a selective growth medium which allows separation of the transformed cells from the parent cells dependent on the nature of the marker sequence of the vector DNA. Preferably, a growth medium is used which does not allow growth of cells which do not contain the vector. The transformation of yeast comprises, for example, steps of enzymatic removal of the yeast cell wall by means of glucosidases, treatment of the obtained spheroplasts with the vector in the presence of polyethylene glycol and Ca²⁺ ions, and regeneration of the cell wall by embedding the spheroplasts into agar. Preferably, the regeneration agar is prepared in a way to allow regeneration and selection of the transformed cells as described above at the same time.

Transformation of cells of higher eucaryotic origin, such as mammalian cell lines, is preferably achieved by transfection. Transfection is carried out by conventional techniques, such as calcium phosphate precipitation, microinjection, protoplast fusion, electroporation, i.e. introduction of DNA by a short electrical pulse which transiently increases the

permeability of the cell membrane, or in the presence of helper compounds such as diethylaminoethyldextran, dimethyl sulfoxide, glycerol or polyethylene glycol, and the like. After the transfection procedure, transfected cells are identified and selected, for example, by cultivation in a selective medium chosen depending on the nature of the selection marker, for example standard culture media such as Dulbecco's modified Eagle medium (DMEM), minimum essential medium, RPMI 1640 medium and the like, containing e.g. the corresponding antibiotic.

The host cells are transformed with the recombinant L-chain gene construct alone, with the recombinant H-chain gene construct alone, with both, either sequentially or simultaneously, or by using a vector construct comprising both the L-chain and H-chain genes, for example a recombinant single-chain antibody gene construct as indicated hereinbefore.

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Preferred are host cells transformed with a recombinant single-chain antibody gene construct comprising DNA coding for the heavy chain variable domain of an anti-c-erbB-2 antibody, DNA coding for a spacer group, DNA coding for the light chain variable domain of an anti-c-erbB-2 antibody and DNA coding for an effector molecule, in particular transfected with the preferred recombinant single-chain antibody gene construct as indicated hereinbefore. Further examples of host cells of the invention are cells transfected with similar recombinant plasmids which contain alternative orientations of the H- and L-chain gene constructs, and those incorporating additional DNA elements to facilitate high levels of expression of the recombinant antibodies.

The host cells of the invention are genetically stable, secrete recombinant antibodies of the invention of constant specificity and can be activated from deep-frozen cultures by thawing and recloning.

The transformed host cells are cultured by methods known in the art in a liquid medium containing assimilable sources of carbon, e.g. carbohydrates such as glucose or lactose, nitrogen, e.g. amino acids, peptides, proteins or their degradation products such as peptones, ammonium salts or the like, and inorganic salts, e.g. sulfates, phosphates and/or carbonates of sodium, potassium, magnesium and calcium. The medium furthermore contains, for example, growth-promoting substances, such as trace elements, for example iron, zinc, manganese and the like.

The medium is preferably so chosen as to exert a selection pressure and prevent the growth of cells which have not been transformed or have lost the hybrid vector. Thus, for example, an antibiotic is added to the medium if the hybrid vector contains an antibiotic resistance gene as marker. If, for instance, a host cell is used which is auxotrophic in an essential amino acid whereas the hybrid vector contains a gene coding for an enzyme which complements the host defect, a minimal medium deficient of said amino acid is used to culture the transformed cells.

Cells of higher eucaryotic origin such as mammalian cells are grown under tissue culture conditions using commercially available media, for example Dulbecco's modified Eagle medium (DMEM), minimum essential medium, RPMI 1640 medium and the like as mentioned above, optionally supplemented with growth-promoting substances and/or mammalian sera. Techniques for cell cultivation under tissue culture condition are well known in the art and include homogeneous suspension culture, e.g. in an airlift reactor or in a continuous stirrer reactor, or immobilized or entrapped cell culture, e.g. in hollow fibres, microcapsules, on agarose microbeads, porous glass beads, ceramic cartridges, or other microcarriers.

Culturing is effected by processes which are known in the art. The culture conditions, such as temperature, pH value of the medium and fermentation time, are chosen so that a maximum titer of the polypeptide or derivative of the invention is obtained. Thus, an <u>E. coli</u> or yeast strain is preferably cultured under aerobic conditions by submerged culture with shaking or stirring at a temperature of about 20°C to 40°C, preferably at about 30°C, and a pH value of 4 to 8, preferably of about pH 7, for about 4 to 30 hours, preferably until maximum yields of the polypeptide or derivative of the invention are reached.

When the cell density has reached a sufficient value, the culture is interrupted and the polypeptide or derivative can be isolated. If the hybrid vector contains a suitable secretion signal sequence, the polypeptide or derivative is secreted by the transformed cell directly into the culture medium. Otherwise, the cells have to be destroyed, for example by treatment with a detergent such as SDS, NP-40TM, TritonTM or deoxycholic acid, lysed with lysozyme or a similarly acting enzyme, or disrupted by an osmotic shock or ultra-sound. Break-up of the cells will also be required if the signal sequence directs the secretion of the desired protein into the cell periplasm. If yeast is used as a host microorganism, the cell wall may be removed by enzymatic digestion with a glucosidase. Alternatively or additionally, mechanical forces, such as shearing forces (e.g. French press, Dyno mill and the like) or shaking with glass beads or aluminium oxide, or alternating freezing, for example in liquid nitrogen, and thawing, for example at 30°C to 40°C, as well as ultra-sound can be used to break the cells.

The cell supernatant or the solution obtained after centrifugation of the mixture obtained after breaking the cells, which contains proteins, nucleic acids and other cell constituents, is enriched in proteins, including the polypeptides of the invention, in a manner which is known <u>per se</u>. Thus, for example, most of the non-protein constituents are removed by polyethyleneimine treatment and the proteins including the polypeptides and derivatives of the invention are precipitated, for example, by saturation of the solution with ammonium sulfate or with other salts. Otherwise, the cell supernatant or lysate is directly pre-purified by filtering through suitable membranes and/or with chromatographic methods, for example affinity chromatography.

The recombinant antibodies and the monoclonal antibodies according to the invention can be used for the qualitative and quantitative determination of the extracellular domain of the growth factor receptor c-erbB-2. This is especially useful for the monitoring of tumor progression, for the decision whether a tumor is amenable to treatment with the recombinant or monoclonal antibodies of the invention, and for monitoring the treatment of tumor with chemotherapy. Tumors considered are those over-expressing c-erbB-2, for example breast and ovarian tumors.

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In general, the monoclonal and the recombinant antibodies according to the invention can be used in any of the known immunoassays which rely on the binding interaction between the antibodies and the antigen, i.e. the extracellular domain of the c-erbB-2 protein. Examples of such assays are radio-, enzyme, fluorescence, chemiluminescence, immunoprecipitation, latex agglutination, and hemagglutination immunoassays, and, in particular, immunostaining methods.

The antibodies according to the invention can be used as such or in the form of enzyme-conjugated derivatives in an enzyme immunoassay. Any of the known modifications of an enzyme immunoassay can be used, for example soluble phase (homogeneous) enzyme immunoassay, solid phase (heterogeneous) enzyme immunoassay, single enzyme immunoassay or double (sandwich) enzyme immunoassay with direct or indirect (competitive) determination of the c-erbB-2 protein.

An example of such an enzyme immunoassay is a sandwich enzyme immunoassay in which a suitable carrier, for example the plastic surface of a microtiter plate or of a test tube, e.g. of polystyrene, polypropylene or polyvinylchloride, glass or plastic beads, filter paper, dextran etc. cellulose acetate or nitrocellulose sheets, magnetic particles or the like, is coated with a monoclonal antibody of the invention by simple adsorption or optionally after activation of the carrier, for example with glutaraldehyde or cyanogen bromide. Then test solutions containing the soluble c-erbB-2 protein and finally single-chain recombinant antibodies of the invention comprising a detectable enzyme, e.g. alkaline phosphatase, are added. The amount of the soluble c-erbB-2 protein in the test solution is directly proportional to the amount of bound recombinant antibody and is determined by adding an enzyme substrate solution. The enzyme substrate reaction results, for example, in a color change which can be observed by eye or with optical measuring devices.

The antibodies according to the invention can be used as such or in the form of radioactively labelled derivatives in a radioimmunoassay (RIA). As described above for enzyme immunoassays, any of the known modifications of a radioimmunoassay can be used.

The tests are carried out in an analogous manner to the enzyme immunoassays described above using a radioactive label, e.g. ¹²⁵l, instead of an enzyme label. The amount of immune complex formed which corresponds to the amount of c-erbB-2 protein present in the test solutions is determined by measuring the radioactivity of the immune complex.

For immunostaining cryosections of cryopreserved biopsy material or paraffin embedded tissue sections are treated with a solution containing a recombinant antibody of the invention comprising a detectable enzyme. Bound recombinant antibody is detected by treatment with a suitable enzyme substrate, preferably an enzyme substrate which leads to a solid deposit (stain) at the site of the recombinant antibody of the invention. In place of recombinant antibodies comprising an enzyme, a recombinant antibody comprising streptavidin and a solution of a biotin-enzyme-conjugate may be used, which leads to higher enzyme concentration at the site of the antibody and hence increased sensitivity of the immunostaining method. The solid deposit of the enzyme substrate is detected by inspection with a microscope, for example with a fluorescence microscope, or by scanning the optical density at the wavelength of the stain.

The use according to the invention of recombinant and/or monoclonal antibodies as described hereinbefore for the determination of c-erbB-2 protein also includes other immunoassays known <u>per se</u>, for example immunofluorescence assays, latex agglutination with antibody-coated or antigen coated latex particles, hemagglutination with antibody-coated or antigen-coated red blood corpuscles, evanescent light assays using an antibody-coated optical fibre and other direct-acting immunosensors which convert the binding event into an electrical or optical signal, or the like.

The invention also concerns test kits for the qualitative and quantitative determination of c-erbB-2 protein comprising recombinant antibodies of the invention and/or monoclonal antibodies of the invention and, optionally, adjuncts.

Test kits according to the invention for an enzyme immunoassay contain, for example, a suitable carrier, optionally treeze-dried solutions of a monoclonal antibody, optionally freeze-dried or concentrated solutions of a recombinant antibody comprising an enzyme or streptavidin, solutions of an enzyme-biotin conjugate if a recombinant antibody comprising streptavidin is used, enzyme substrate in solid or dissolved form, standard solutions of c-erbB-2 protein, buffer solutions, and, optionally, polypeptides or detergents for preventing non-specific adsorption and aggregate formation, pipettes, reaction vessels, calibration curves, instruction manuals and the like.

Test kits according to the invention for immunostaining contain, for example, optionally freeze-dried or concentrated solutions of a recombinant antibody comprising an enzyme or streptavidin, solutions of an enzyme-biotin conjugate if a recombinant antibody comprising streptavidin is used, enzyme substrate in solid or dissolved form, buffer solutions, and, optionally, pipettes, reaction vessels, calibration curves, instruction manuals and the like.

The recombinant and monoclonal antibodies of the invention can be used for the qualitative and quantitative determination of c-erbB-2 protein. Due to the fact that the growth factor receptor c-erbB-2 is overexpressed in certain tumor types, for example breast and ovarian tumors, the antibodies are particularly well suited for detection and monitoring of the mentioned tumors. In addition, radiolabelled derivatives of the antibodies of the invention may be used for the in vivo localization of tumors in a patient using radioscanning techniques. To that end, radiolabelled derivatives of antibodies of the invention are injected into the patient, and the patient scanned with a gamma imager at regular intervals. Cells over-expressing the growth factor receptor c-erbB-2 will take up more radioactive antibodies than other tissue and will be clearly recognized by the gamma imaging camera. Preferentially recombinant or monoclonal antibodies labelled with ¹³¹I or with ^{99m}Tc are used for radioscanning in amounts of 3 to 8 μg representing 15 to 30 μCi per kg body weight.

The antibodies of the invention can further be used for the isolation and purification of the c-erbB-2 protein from natural sources or from transformed host cells by immunoaffinity chromatography.

Furthermore, the monoclonal antibodies and the recombinant antibodies of the invention, in particular recombinant antibodies comprising an effector molecule, especially a toxin, in particular Pseudomonas exotoxin, are useful for the treatment of patients with tumors over-expressing the growth factor receptor c-erbB-2, for example breast or ovarian tumors. If it is desired, tumor therapy may comprise applying more than one, e.g. two different, antibodies of the invention, for example applying both FRPS and FWP51. The recombinant antibodies comprising a phosphatase may be used in connection with a phosphorylated prodrug such as mitomycin phosphate or etoposide phosphate, thus enabling the conversion of the active drug to the prodrug at the site of the tumor.

The invention therefore also concerns pharmaceutical compositions for treating tumors over-expressing the growth factor receptor c-erbB-2 comprising a therapeutically effective amount of a recombinant antibody or of a monoclonal antibody according to the invention and a pharmaceutically acceptable carrier. Preferred are pharmaceutical compositions for parenteral application. Compositions for intramuscular, subcutaneous or intravenous application are e.g. isotonic aqueous solutions or suspensions, optionally prepared shortly before use from lyophilized or concentrated preparations. Suspensions in oil contain as oily component the vegetable, synthetic or semi-synthetic oils customary for injection purposes. The pharmaceutical compositions may be sterilized and contain adjuncts, e.g. for conserving, stabilizing, wetting, emulsifying or solubilizing the ingredients, salts for the regulation of the osmotic pressure, buffer and/or compounds regulating the viscosity, e.g. sodium carboxycellulose, carboxymethylcellulose, sodium carboxymethylcellulose, dextran, polyvinylpyrrolidine or gelatine.

The pharmaceutical compositions of the invention contain from approximately 0.01% to approximately 50% of active ingredients. They may be in dosage unit form, such as ready-to-use ampoules or vials, or also in lyophylized solid form.

In general, the therapeutically effective dose for mammals is between approximately 5 and 25 µg of a recombinant antibody of the invention or of a monoclonal antibody of the invention per kg body weight depending on the type of antibody, the status of the patient and the mode of application. The specific mode of administration and the appropriate dosage will be selected by the attending physician taking into account the particulars of the patient, the state of the disease, the type of tumor treated, and the like. The pharmaceutical compositions of the invention are prepared by methods known in the art, e.g. by conventional mixing, dissolving, confectioning or lyophilizing processes. Pharmaceutical compositions for injection are processed, filled into ampoules or vials, and sealed under aseptic conditions according to methods known in the art.

The invention particularly concerns the monoclonal antibodies, the hybridoma cell lines, the recombinant singlechain antibodies, the recombinant DNAs, the transformed host cells, and the methods for the preparation thereof as described in the Examples. The following examples illustrate the invention but do not limit it to any extent.

Abbreviations

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45	ATP	adenosine triphosphate
	BSS	Earle's balanced salt solution
	BSA	bovine serum albumin
	DEAE	diethylaminoethyl
	DMEM	Dulhecco's modified Facile's med

DMEM Dulbecco's modified Eagle's mediur dNTP deoxynucleotide triphosphate

DTT dithiothreitol

EDTA disodium ethylenediaminetetraacetate

EGF epidermal growth factor

EGTA ethyleneglycol-bis-(β-aminoethyl ether)-N,N,N',N'-tetraacetic acid

55 FCS fetal calf serum

HAT medium hypoxanthine, aminopterin and thymidine medium

HEPES N-2-hydroxyethylpiperazine-N'-2-ethanesulfonic -2-ethanesulfonic acid

HT medium hypoxanthine and thymidine medium

lg immunoglobulin

IPTG isopropyl-β-thiogalactoside
MAb monoclonal antibody
PBS phosphate-buffered saline
PCR polymerase chain reaction
PMSF phenylmethylsulfonyl fluoride

SDS-PAGE sodium dodecyl sulfate-polyacrylamide gel electrophoresis

Tris Tris-(hydroxymethyl)-aminomethane

U unit

V_L light chain variable domain V_H heavy chain variable domain

XP 5-bromo-4-chloro-3-indolyl phosphate p-toluidine salt

Examples

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Example 1. Preparation of hybridoma cell lines FRPS, FSP16, FWP51 and FSP77

- 1.1 Source of antigen and immunization of Balb/c mice: The SKBR3 human breast tumor cell line (ATCC HTB 30), isolated in 1970 from a pleural effusion of a breast cancer patient, expresses approximately 1 x 10⁶ molecules of the c-erbB-2 receptor protein per cell. 20 x 10⁶ SKBR3 cells in PBS are injected subcutaneously and/or intraperitoneally into Balb/c mice. The cells are mixed 1:1 (v/v) with complete Freund's adjuvant. The injections are repeated a total of five times over the period of approximately 3 months replacing Freund's incomplete adjuvant for complete adjuvant. The final injection of cells is given three days before the fusion.
- 1.2 <u>Cell fusion:</u> Immunized mice are sacrificed and their splenocytes fused according to conventional methods (Koehler & Milstein, Nature 256: 495, 1976). Spleen cells are mixed at a 5:1 to 10:1 ratio with the fusion partner, the mouse myeloma cell line PAI (Stoker et al., Research Disclosure #21713, 1982), in the presence of 41 % polyethylene glycol 4000 (Merck). Fused cells are plated at a density of 1 x 10⁶ cells per well in 24-well microtiter plates on peritoneal macrophages and fed 3 times per week with standard HAT selection medium for 2 weeks followed by 2 weeks of HT medium. When the growth of hybridoma cells becomes visible, the supematants are screened as described in Example 1.3. Positive hybridomas are cloned and stored.
- 1.3 Antibody detection in hybridoma supernatants: Culture fluids of growing hybridomas are tested for the presence of anti-c-erbB-2 antibody using a protocol involving two steps, immunofluorescence and immunoprecipitation.
- 1.3.1 Immunofluorescence: In the first step, hybridoma supernatants are tested for their immunofluorescent staining of mouse cells expressing high levels of the human c-erbB-2 protein. To isolate these cells the HC11 mouse mammary epithelial cell line (Ball et al., EMBO J. 7: 2089, 1988) is transfected according to conventional, previously described methods (Graham & van der Eb, Virology 52: 456, 1973) with a plasmid expressing the human c-erbB-2 protein (Masuko et al., Jpn. Cancer Res. 80: 10, 1989) and with the plasmid pSV2neo (Southem & Berg, J. Mol. Appl. Genet. 1: 327, 1982) which encodes the gene for resistance to the drug G418. Transfected cells are selected 2 weeks in medium containing 200 µg/ml G418 (Geneticin, Gibco-BRL). Individual clones are selected and analyzed for expression of the human c-erbB-2 protein using conventional protein blotting techniques (Towbin et al., Proc. Natl. Acad. Sci. USA 76: 4350, 1979). A clone expressing high levels of the human c-erbB-2 protein (clone R1#11) is selected and used in the immunofluorescent assay. Non-transfected HC11 cells serve as control cells.

The assay is done in the following manner: The cells (R1 #11 or HC11) are grown in RPMI medium containing 8 % heat inactivated FCS (Amimed), 10 ng/ml EGF (Inotech) and 5 µg/ml insulin (Sigma) for 1 -2 days on fibronectin (Boehringer Mannheim) coated cover slips. Fibronectin coated cover slips are prepared and stored at room temperature and they are used routinely for screening. The coverslips are rinsed in PBS containing calcium and magnesium and fixed by treatment for 10 min with 3.7 % formaldehyde (v/v in PBS). To reduce the non-specific binding the coverslips are incubated 20 min in PBS containing 3 % BSA (Sigma). The coverslips are washed in PBS and in water, then allowed to dry at room temperature. 20 - 30 µl of hybridoma supernatants are added to circled areas on a coverslip which is incubated 1 - 2 h at room temperature in a humified atmosphere. The coverslips are then washed three times with PBS containing 0.05 % Triton-X100TM (Fluka) and incubated an additional hour with anti-mouse Ig, fluorescein-linked whole antibody from sheep (Amersham). After three washes with PBS and one wash with water the cells are screened for fluorescence using a fluorescence microscope and a water immersion lens. Those hybridoma supernatants which are positive are screened in the second step described in Example 1.3.2.

1.3.2 Immunoprecipitation and protein blotting analysis: The SKBR3 human breast tumor cells express approximately 1 x 10⁶ molecules of the c-erbB-2 protein per cell. A cell lysate is prepared by extracting approximately 4 x 10⁶ cells in 1 ml of buffer containing 1 % Triton-X100™ (Fluka), 50 mM Tris-HCl, pH 7.5, 5 mM EGTA, 0.15 M NaCl, 1 mM PMSF (Boehringer Mannheim), 80 µg/ml aprotinin (Boehringer Mannheim), 50 µg/ml leupeptin (Boehringer Mannheim),

and 4 µg/ml pepstatin (Boehringer Mannheim). 200 -500 µl supernatant of hybridomas which are positive in the immunofluorescence assay described in Example 1.3.1 are incubated with 100 µl of the SKBR3 extract (2.5 - 4.0 mg/ ml). This amount of extract contains approximately 50 - 100 ng of c-erbB-2 protein. The hybridoma supernatants and SKBR3 extract are incubated overnight on ice, then 1 µl of the IgG fraction of sheep anti-mouse Ig (ICN Immunobiologicals) is added. The complexes are collected by the addition of Protein-A Sepharose™ (Pharmacia), washed with TNET (140 mM NaCl, 50 mM Tris-HCl, pH 7.5, 5 mM EDTA, 1 % Triton X-100™) and water, boiled in sample buffer (80 mM Tris-HCl, pH 6.8, 0.2 % SDS, 10 % glycerol) and the supernatants loaded onto 8 % SDS-PAGE. The proteins are electrophoresed and blotted onto PVDF membranes (Millipore) using a technique originally described by Towbin et al. (Proc. Natl. Acad. Sci. USA 76: 4350, 1979) with some modifications. The proteins are transferred using a semidry blotter (G. Frobel, Model 1004.01) following the instructions of the manufacturer. The membranes are blocked in PBS containing 0.5 % gelatin (Merck) for 1 h at 37°C. The membranes are washed twice for 5 min at 37°C in PTG (PBS containing 0.02 % gelatin (Merck) and 0.25 % Triton-X100™ (Fluka)). The c-erbB-2 protein is detected by incubating the membrane 45 min at 37°C in PTG containing an antiserum which is raised against the carboxy terminal 13 amino acids of the c-erbB-2 protein (Gullick et al., Int. J. Cancer 40: 246, 1987, antiserum 21N). The membranes are washed 3 times for 5 min at 37°C in PTG. The membrane-bound 21N antiserum is detected by incubating the membrane in PTG containing 0.1 µC/ml 125I-labeled protein-A (Amersham). The membranes are washed 4 times for 5 min at 37°C in PTG and exposed to X-ray film. The hybridomas whose supernatants are able to specifically immunoprecipitate the c-erbB-2 protein are grown for single cell cloning and further characterization described below.

Example 2. Characterization of c-erbB-2 specific MAbs

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- 2.1 <u>Hybridoma storage and processing</u>: Hybridoma FRP5, FSP16, FWP51 and FSP77 secreting anti-c-erbB-2 MAb FRP5, FSP16, FWP51 and FSP77, respectively, can be grown in culture, frozen at -80°C or in liquid nitrogen and recultivated. The cells are cloned by the method of limiting dilution and have been deposited with the European Collection of Animal Cell Lines in England. The hybridoma cell lines have the following access numbers: FRP5: 90112115, FSP16: 90112116, FSP77: 90112117, FWP51: 90112118. The cells are expanded by forming ascites in Balb/c mice primed with pristane. The antibodies are purified from the ascites by ammonium sulfate precipitation and ion exchange chromatography on DE 52 DEAE-cellulose columns (Whatman). Purified MAbs are stored in PBS at -80°C.
- 2.2 <u>Isotyping of the MAbs</u>: The isotype of the MAbs FRP5, FSP16, FWP51 and FSP77 is determined by ELISA analysis with rabbit antisera to mouse Ig classes and sub-classes (Biorad Mouse Typer TMSub Isotyping KitTM) as per manufacturer's suggested procedure. MAbs FRP5, FWP51, and FSP77 are of the IgG1 isotype, while FSP16 is of the IgG2b isotype. The light chains of all the MAbs are of the kappa type.
- 2.3 Flow cytometry: A FACS analysis using the c-erbB-2 specific MAbs is carried out as follows: SKBR3 human breast tumor cells are trypsinized, washed in FACS medium (BSS containing 10 μM sodium azide, 4 % FCS and 25 mM EDTA), and 1 x 10⁶ cells are resuspended in 100 μl of FACS medium. Non-specific binding sites are blocked by incubating the cells 10 min at room temperature with 5 μl of goat serum. The SKBR3 cells are collected by centrifugation, resuspended in 50 μl of a 1:2 dilution of the supernatant made in FACS medium and incubated 45 min on ice. The cells are washed with 4 ml FACS medium, collected by centrifugation, resuspended in 50 μl of FACS medium containing a 1:20 dilution of anti-mouse lg, fluorescein-linked whole antibody from sheep (Amersham), and incubated for 30 min on ice. 4 ml of FACS medium are added, the cells are collected by centrifugation, resuspended in 100 μl of FACS medium and analyzed without fixation for their fluorescence in a Becton-Dickinson FACScanTM. As a control, SKBR3 cells are incubated with a non-reacting IgG1 MAb (1236S31-3). The FACS analysis shows that the SKBR3 cells treated with MAb FRPS, FSP16, FWP51, and FSP77 have a higher fluorescence than cells treated with the control MAb. These results show that the MAbs bind to the extracellular domain of the c-erbB-2 protein.
- 2.4 <u>Binding domain of c-erbB-2 specific MAbs.</u> MAbs FRPS and FSP77 are covalently linked with ¹²⁵I (as carrier free sodium ¹²⁶iodide, Amersham) to a specific activity of 1 µC/µg using lodogen (1,3,4,6-tetrachloro-3a,6a-diphenylg-lycouril, Sigma) according to a standard protocol (Antibodies: A Laboratory Manual, Cold Spring Harbor Laboratory, 1988, p. 330). Competition experiments are conducted by incubating SKBR3 cells (0.5 1 x 10⁵ cells per 15 mm well, Nundon™ 4-well multidish) with 250 µl RIA buffer (120 mM NaCl, 50 mM HEPES, pH 7.8, 1 mM EDTA, 2 % BSA) containing labeled FRPS or FSP77 and varying amounts of unlabeled MAb FRPS, FSP16, FWP51 and FSP77 for 2 h at 4°C. The cells are washed 5 times with the RIA buffer, solubilized in 0.5 ml 1 % Triton X-100™, 10 % glycerol, 20 mM HEPES, pH 7.4, for 30 min at room temperature and the bound radioactivity is measured in a gamma counter. The results show that MAbs FRPS and FSP16 compete with each other for binding to SKBR3 cells which suggests that these 2 MAbs bind to the same domain on the c-erbB-2 protein. MAbs FWP51 and FSP77 neither compete with each other nor with FRPS or FSP16 for binding to the c-erbB-2 protein. In conclusion, the panel of 4 MAbs bind to 3 different domains of the extracellular portion of the c-erbB-2 membrane receptor tyrosine kinase.

Example 3. Isolation of RNA from the hybridoma cell line FRP5

- 3.1 <u>Growth of FRPS cells:</u> FRPS hybridoma cells (1 x 10⁸) are grown in suspension culture at 37°C in DMEM (Seromed) further containing 10 % FCS (Amimed), 1 mM sodium pyruvate (Seromed), 2 mM glutamine (S romed), 50 μM 2-mercaptoethanol and 100 μg/ml of gentamycin (Seromed) in a humidified atmosphere of air and 7.5 % CO₂ in 175 cm tissue culture flasks (Falcon 3028). The cells are harvested by centrifugation, washed once in PBS, flash frozen in liquid nitrogen and kept frozen as a pellet at 80°C in a clean, sterile plastic capped tube.
- 3.2 Extraction of total cellular RNA from FRP5 cells: Total RNA is extracted using the acid guanidinium thiocyanate-phenol-chloroform method described by Chomczynski & Sacchi (Anal. Biochem. 162: 156, 1987). Cell pellets of FRPS cells (1 x 10⁸) are thawed directly in the tube in the presence of 10 ml of denaturing solution (4 M guanidinium thiocyanate (Fluka), 25 mM sodium citrate, pH 7.0, 0.5 % N-lauroylsarcosine (Sigma), 0.1 M 2-mercaptoethanol). The solution is homogenized at room temperature. Sequentially, 1 ml of 2 M sodium acetate, pH 4, 10 ml of phenol (water saturated) and 2 ml of chloroform-isoamyl alcohol mixture (49:1) are added to the homogenate. The final suspension is shaken vigorously for 10 sec and cooled on ice for 15 min. The samples are centrifuged at 10,000 x g for 20 min at 4°C. After centrifugation, RNA which is present in the aqueous phase is mixed with 10 ml of isopropanol and placed at -20°C for 1 h. The RNA precipitate is collected by centrifugation, the pellet dissolved in 3 ml water and the RNA reprecipitated by addition of 1 volume of isopropanol at -20°C. After centrifugation and washing the pellet in ethanol, the final pellet of RNA is dissolved in water. The method yields approximately 300 µg of total cellular RNA. The final purified material is stored frozen at -20°C.
- 3.3 <u>Isolation of poly(A) containing RNA:</u> Poly(A) containing RNA is selected from total RNA by chromatography on oligo(dT)-cellulose (Boehringer Mannheim) as described originally by Edmonds et al. (Proc. Natl. Acad. Sci. USA 68: 1336, 1971) and modified by Maniatis et al. (Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory, 1982, p. 197). The poly(A)-containing RNA is prepared as described in the published procedure with the exception that the RNA is eluted from the oligo(dT)-cellulose with water rather than SDS-containing buffer. The poly(A)-containing RNA is precipitated with ethanol and collected by centrifugation. The yield of poly(A)-containing RNA is approximately 30 μg from 300 μg of total cellular RNA. The final purified material is stored frozen at -20°C.

Example 4. Cloning of functional heavy and light chain rearrangements from the FRP5 hybridoma cell line

Poly(A)-containing RNA isolated from FRP5 hybridoma cells as described in Example 3.3 provides the source for cDNA synthesis and subsequent amplification of V-region minigenes. Amplification products of the expected size are purified from agarose gels and cloned into appropriate vectors. Functional rearrangements are identified by sequencing.

4.1 Oligonucleotides:

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MCK2 is designed to be complementary to a region in the murine immunoglobulin κ (kappa) constant minigene.

5' - TCACTGGATGGTGGGAAGATGGA - 3'

MCHC2 is designed to be complementary to a region in the murine immunoglobulin γ1 constant minigene.

5' - AGATCCAGGGGCCAGTGGATAGA - 3'

The oligonucleotides VH1FOR, VH1BACK, VK1FOR, and VK1BACK are designed by Orlandi et al. (Proc. Natl. Acad. Sci. USA 86: 3833, 1989) to match consensus sequences.

VH1FOR: 5'-TGAGGAGACGGTGACCGTGGTCCCTTGGCCCCAG-3'

VH1BACK: 5' - AGGT(C/G)(C/A)A(G/A)CTGCAG(G/C)AGTC(T/A)GG - 3'

VK1FOR: 5' - GTTAGATCTCCAGCTTGGT(C/G)C(C/G) - 3'

VK1BACK: 5' - GACATTCAGCTGACCCAGTCTCCA - 3'

4.2 <u>cDNA synthesis</u>: 55 ng of poly(A)-containing RNA is dissolved in a buffer containing 50 mM Tris-HCl, pH 8.3, 3 mM magnesium chloride, 10 mM DTT, 75 mM KCl, 400 μM dNTPs (N = G, A, T and C), 100 μg BSA (molecular biology grade, Boehringer Mannheim), 100 U RNAse inhibitor (Boehringer Mannheim), 25 pmol MCK2 and 25 pmol MCHC2. The RNA is denatured at 70°C for 5 min and then chilled on ice for 2 min. After addition of 200 U of MMLV reverse transcriptase (Gibco, BRL) cDNA synthesis is achieved by incubation for 1 h at 37°C.

4.3 <u>Polymerase chain reaction:</u> One tenth of the cDNA reaction is used for DNA amplification in buffer containing 10 mM Tris-HCl, pH 8.3, 1.5 mM MgCl₂, 50 mM KCl, 10 mM β-mercaptoethanol, 200 μM dNTPs (N= G, A, T and C), 0.05 % Tween-20™ (Merck), 0.05 % NP-40™ (Merck), 10 % DMSO (Merck), 25 pmol oligonucleotide 1 (see below), 25 pmol oligonucleotide 2 (see below) and 2.5 U Amplitaq™ DNA polymerase (Perkin Etmer Cetus). Taq polymerase is added after initial denaturation at 93°C for 1 min and subsequent annealing at 37°C. In the first 4 cycles primer extension is performed at 71°C for 0.2 min, denaturation at 93°C for 0.01 min and annealing at 37°C for 0.2 min. For the last 25 cycles the annealing temperature is raised to 62°C. Finally, amplification is completed by a 3 min primer extension step at 71°C.

PCR Product	oligonucleotide 1	oligonucleotide 2
HC	MCHC2	VH1BACK
н	VH1FOR	VH1BACK
LC	MCK2	VH1BACK
L	VK1FOR	VH1BACK

4.4 <u>Modification and purification</u>: Amplified material is extracted with CHCl₃ and precipitated with ethanol in the presence of 200 mM LiCl. To facilitate cloning, blunt ends are created by a 3 min treatment with 1 U T4 DNA polymerase (Boehringer Mannheim) in 66 mM Tris-acetate, pH 7.9, 132 mM potassium acetate, 20 mM magnesium acetate, 1 mM DTT, 200 μg/ml BSA (molecular biology grade, Boehringer Mannheim), and 400 μM dNTPs (N = G, A, T and C). The polymerase is inactivated by heating for 15 min at 65°C before phosphorylation of the DNA with 10 U T4 polynucleotide kinase (Pharmacia) at 37°C for 1 h. For this purpose the buffer is adjusted to 50 mM EDTA and 1 mM ATP. The modified amplification products are separated on a 1.2 % (w/v) agarose gel (ultra pure DNA grade agarose, Biorad) and DNA of the expected size is eluted by means of DEAE NA 45 membranes (Schleicher & Schuell).

4.5 <u>Ligation</u>: Bluescript™ KS+ (70 ng) linearized with Xbal, treated with Klenow DNA polymerase (Boehringer Mannheim) to give blunt ends and dephosphorylated with calf intestinal phosphatase, and 30 ng of purified amplification product are ligated using 0.5 U T4 DNA ligase (New England Biolabs) in 50 mM Tris-HCl, pH 7.8, 10 mM magnesium chloride, 10 mM DTT, and 0.8 mM ATP overnight at 16°C. One half of the ligation mixture is used to transform <u>E. coli</u> K803 to obtain ampicillin resistant colonies. These are screened for the desired ligation products using a NaOH based plasmid "miniprep" method (Maniatis et al., Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory, 1982). The following plasmids are obtained:

PCR product	Plasmid clones
HC	pMZ15/1
	pMZ15/2
н	pMZ16/1
	pMZ16/2
L	pMZ17/1
	pMZ17/2
LC	pMZ18/1
	pMZ18/2

4.6 <u>Sequencing</u>: Sequencing is done using Sequenase™ kits (United States Biochemicals) with T3 and T7 oligonucleotide primers according to procedures provided by the manufacturer.

Plasmid pMZ17/1 contains a non-functional rearrangement. Plasmid pMZ17/2 contains an Ig-unrelated sequence. Plasmids pMZ18/1 (SEQ ID NO:2) and pMZ18/2 contain identical functional FRPS kappa light chain variable domain inserts. Plasmids pMZ16/1 (SEQ ID NO:1) and pMZ16/2 contain identical functional FRPS heavy chain variable domain inserts. Plasmids pMZ15/1 and pMZ15/2 also contain FRPS heavy chain variable domain inserts together with some constant region DNA. Plasmids pMZ16/1 and pMZ18/1 are used as a source for further subcloning steps.

Example 5. Construction of the MAb FRP5 single-chain Fv gene

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5.1 Construction and sequence of a cloning linker for the heavy and light chain variable domain cDNAs: Using oligonucleotides, a linker sequence which allows the cloning of PCR amplified mouse heavy chain variable domain cDNA as a Pstl/BstEll fragment and of PCR amplified mouse kappa light chain variable domain cDNA as a Pvull/Bglll fragment is constructed. This creates an open reading frame in which heavy and light chain variable domains are

For the construction of the cloning linker the 6 complementary oligonucleotides 1A, 1B, 2A, 2B, 3A, 3B are used.

- 1A: 5'-CAAGCTTCTCAGGTACAACTGCAGGAGGTCACCGTTTCCTCTGGCGG-3'
- 1B: 5'-GAAACGGTGACCTCCTGCAGTTGTACCTGAGAAGCTTGCATG-3'
- 2A: 5'-TGGCGGTTCTGGTGGCGGTGGCTCCGGCGGTGGCGGTTCTGAC-3'
- 2B: 5'-GCCACCGCCGGAGCCACCGCCACCAGAACCGCCACCGCCAGAG-3'
- 3A: 5'-ATCCAGCTGGAGATCTAGCTGATCAAAGCT-3'

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3B: 5'-CTAGAGCTTTGATCAGCTAGATCTCCAGCTGGATGTCAGAACC-3'

40 pM of oligonucleotides 1B, 2A, 2B, 3A are phosphorylated at the 5' end using T4 polynucleotide kinase (Boehringer Mannheim) in four separate reactions in a total volume of 20 µl following the method described by Maniatis et al. (Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory, 1982). Oligonucleotides 1A and 3B are not phosphorylated in order to avoid self ligation of the linker in the final ligation reaction. After the kinase reaction, the enzyme is inactivated by incubation at 70°C for 30 min. In three separate reactions, each containing 40 pM of two oligonucleotides in a total volume of 40 µl, non-phosphorylated 1A and phosphorylated 1B, phosphorylated 2A and phosphorylated 2B, and phosphorylated 3A and non-phosphorylated 3B are mixed. Hybridization of the oligonucleotides in the three reactions is carried out by heating to 95°C for 5 min, incubation at 65°C for 5 min and slowly cooling to room temperature. 10 µl from each of the three reactions are mixed, 4 µl of 10 x ligation buffer (Boehringer) and 4 units of T4 DNA ligase (Boehringer) are added and the total volume is adjusted to 40 µJ with sterile water. The annealed pairs of oligonucleotides are ligated into one linker sequence for 16 h at 14°C. The reaction mixture is extracted with an equal volume of phenol/chloroform (1:1) followed by re-extraction of the aqueous phase with an equal volume of chloroform/isoamylalcohol (24:1). The aqueous phase is collected, 0.1 volumes of 3 M sodium acetate pH 4.8 and 2 volumes of ethanol are added, and the DNA is precipitated at -70°C for 4 h and collected by centrifugation. The resulting linker sequence has a Sphl and a Xbal adaptor end. It is ligated to Sphl and Xbal digested pUC19 in a reaction containing 100 ng of ligated linker and 200 ng of Sphl/Xbal digested pUC19. After transformation into E. coli XL1 Blue™ (Stratagene), plasmid DNA from 4 independent colonies is isolated by the alkaline lysis mini-preparations method (Maniatis et al., Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory, 1982). The DNA sequence of the linker cloned in pUC19 is determined by sequencing double stranded DNA in both directions with Sequenase II (United States Biochemicals) and pUC universal and reverse primers (Boehringer) following the manufacturer's protocol. Three out of the four recombinant pUC19 isolates sequenced contain the correct linker sequence. One of them is designated pWW19 and used in the further experiments. The sequence is shown in SEQ ID NO:3.

- 5.2 <u>Preparation of a plasmid for the subcloning of variable domains:</u> The Fv cloning linker sequence is derived as a 144 bp HindIII/SacI fragment from pWW19 and inserted into HindIII/SacI digested Bluescript™ KS+ (ex PvuII) (Stratagene) which contains no PvuII restriction sites. The resulting plasmid, pWW15, allows cloning of heavy and light chain variable domains as PstI/BstEII and PvuII/BglII fragments, respectively.
- 5.2.1 <u>Subcloning of the FRPS heavy chain variable domain</u>: Plasmid pMZ16/1 is digested with PstI and BstEII and the 338 bp heavy chain variable domain fragment of FRP5 is isolated. It is cloned into PstI/BstEII digested pWW19 yielding the plasmid pWW31.
- 5.2.2 Mutation of the FRPS light chain variable domain and assembly of the Fv fusion gene: To facilitate subcloning of the FRPS light chain variable domain into the Fv cloning linker, a Pvull restriction site and a Bglll restriction site are introduced at the 5' and 3' ends, respectively, of the coding region. The FRPS light chain variable domain coding region is isolated as a Sacl/BamHI fragment from pMZ18/1. Sacl and BamHI are restriction sites of the Bluescript™ polylinker present in pMZ18/1. The fragment contains the complete light chain variable domain fragment of 392 bp amplified by PCR using the oligonucleotide MCK2 (see above). This fragment is mutated and amplified by PCR using the oligonucleotides

V_L5': 5'-GACATTCAGCTGACCCAG-3' and V₁3': 5'-GCCCGTTAGATCTCCAATTTTGTCCCCGAG-3'

for the introduction of a Pvull restriction site at the 5' end (V_L5') and a Bglll restriction site at the 3' end (V_L5') of the kappa light chain variable domain DNA. 20 ng of the FRP5 variable light chain Sacl/BamHI fragment are used as a template in a 100 µl reaction following the PCR conditions described in Example 4.3. The amplified and mutated fragment is isolated after Pvull/Bglll digestion as a 309 bp fragment from a 1.5 % agarose gel and cloned into Pvull/Bglll

digested pWW15 generating plasmid pWW41. The FRPS kappa light chain variable domain is isolated as a BstEll/Xbal fragment from pWW41 and inserted into BstEll/Xbal digested pWW31. Thus the FRPS heavy chain variable domain in pWW31 and the FRPS kappa light chain variable domain are fused to one open reading frame. Double stranded DNA of three independent clones is sequenced with Sequenase II™ kit (United Biochemicals) in both orientations using pUC universal and reverse primers (Boehringer) following the manufacturer's protocol. One of the plasmids carrying the FRPS heavy chain variable domain fused to the mutated FRPS light chain variable domain is selected and designated pWW52. The sequence of the HindIII/Xbal insert in plasmid pWW52 is shown in SEQ ID NO:4.

Example 6. Construction of a single-chain Fv-phosphatase fusion gene expression plasmid

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The MAb FRPS single-chain Fv gene is fused to the bacterial alkaline phosphatase. This chimeric gene encodes a bifunctional molecule which retains binding activity to the c-erbB-2 protein and has enzymatic activity.

6.1 <u>Mutation of the single-chain Fv(FRPS) gene:</u> To allow gene fusion between the single-chain Fv(FRP5) encoding gene from pWW52 and the alkaline phosphatase gene phoA the stop codon at sequence position 729 to 731 in pWW52 (see Example 5.2.3) is deleted as follows: Plasmid DNA of pWW52 is digested with BstEII and BgIII and the linker sequence and FRPS light chain variable domain encoding fragment is isolated. In another digestion, pWW52 is cleaved with BStEII and BcII. Thus, the large fragment containing vector sequences and the FRPS heavy chain variable domain encoding sequence is isolated. The BstEII/BgIII V_L fragment is now inserted into BstEII/BcII cleaved pWW52 containing V_H. In the resulting plasmid, pWW53, the BgIII/BcII junction is determined by sequencing double stranded DNA as described above.

Sequence of the Bglll/Bcll junction in pWW53 (position numbers correspond to position numbers of the HindIII/ Xbal insert in plasmid pWW52, SEQ ID NO:4):

BglII/BclI

ACA AAA TTG GAG ATC AAA GCT CTA GA

714 -728 | 738 - 748

6.2 <u>Mutation of the E.coli alkaline phosphatase gene phoA</u>: For the construction of the Fv(FRP5)-phoA fusion gene the <u>E. coli</u> alkaline phosphatase gene phoA is mutated to generate a Xbal cleavage site in the coding region of phoA near the N terminus of the mature protein and a Sact cleavage site in the 3' untranslated region of phoA. This step facilitates the cloning of the mutated fragment. A pBR322 derivative carrying the recombinant transposon TnPhoA (Manoil & Beckwith, Proc. Natl. Acad. Sci. USA 82: 8129, 1985) is linearized by BgIII cleavage. 20 ng of the linearized template DNA is used for a 100 μl PCR reaction carried out as described previously using oligonucleotides PhoA5' and PhoA3' as primers 1 and 2.

PhoA5': 5'-CCCTCTAGAGCCTGTTCTGGAAAAC-3' PhoA3': 5'-CCCGAGCTCTGCCATTAAG-3'

Following Xbal/SacI digestion of the PCR products, a 1419 bp fragment is isolated from a 1.5 % agarose gel and inserted into Xbal/SacI digested plasmid pUC19. Ligation is carried out as described above. Ligated DNA is transformed into E. coli XL1 Blue™ (Stratagene). Thus, the open reading frame of the mutated phoA gene is fused in frame to the lacZ open reading frame of pUC19. To show that the mutated phoA gene expresses functional alkaline phosphatase, recombinant clones are plated onto LB agar plates containing 100 μg/ml ampicillin, 0.5 mM IPTG (Sigma), and 40 μg/ml XP (Boehringer). Following induction of the lac promoter of pUC 19, a lacZ-phoA fusion protein is expressed. The phosphatase activity of this fusion protein converts the indicator XP to a blue dye. One of the blue colonies is isolated and the presence of the introduced restriction sites is confirmed by digestion of miniprep DNA with Xbal and SacI. Partial 5' and 3' DNA sequences of the mutated phoA gene are obtained by sequencing double stranded DNA as described above. The DNA sequences are included in the assembly of the final Fv(FRP5)-phoA fusion gene sequence shown in SEQ ID NO:5. The isolated plasmid is designated pWW61 and used for further subcloning steps.

6.3 <u>Construction of a FRP5 Fv-phoA expression plasmid</u>: From plasmid pWW19 (see Example 5.1.2) the cloning linker sequence is isolated as a HindIII/EcoRI fragment and inserted into HindIII/EcoRI digested plasmid pINIII-ompA-Hind (Rentier-Delrue et al., Nucl. Acids Res. 16: 8726, 1988) leading to plasmid pWW16.

From pWW61 (see Example 6.2) the mutated phoA gene is isolated as a Xbal/Sacl fragment and inserted into Xbal/Sacl digested pWW53. The resulting plasmid, pWW615, carries the Fv(FRP5) gene fused in frame to the mutated alkaline phosphatase gene. The Fv(FRP5)-phoA gene is isolated as a HindIII/Sacl fragment from pWW615 and inserted into HindIII/Sacl digested plasmid pWW16. This leads to the production of the Fv(FRP5)-phoA expression plasmid pWW616 (see below). All ligations are carried out as described above. Recombinant plasmids are transformed into <u>E</u>.

coli XL1 Blue™ (Stratagene). The constructs are confirmed by restriction enzyme analysis of plasmid DNA isolated by an alkaline mini preparation method (Maniatis et al., Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory, 1982).

In this construct the Fv single-chain antibody of FRP5, genetically fused to the alkaline phosphatase phoA, can be expressed in <u>E</u>. <u>coli</u> following induction with IPTG. The recombinant protein carries the <u>E</u>. <u>coli</u> outer membrane protein A (ompA) signal sequence at the N terminus (encoded by the pINIII-ompA-Hind vector) to facilitate secretion of the protein into the periplasmic space of <u>E</u>. <u>coli</u> expressor cells.

The sequence of the Fv(FRP5)-phoA fusion gene in expression plasmid pWW616 is shown in SEQ ID NO:5. Part of the phoA sequence is assembled from Chang et al., Gene 44: 121, 1986.

Example 7. Expression of Fv(FRP5)-phoA in E. coli

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Plasmid pWW616 is transformed into the phoA negative <u>E</u>. <u>coli</u> strain CC118 (Manoil & Beckwith, Proc. Natl. Acad. Sci. USA 82: 8129, 1985). A recombinant single colony is grown ovemight in 50 ml LB medium containing 70 μg/ml ampicillin. The ovemight culture is diluted 1:10 in 500 ml fresh LB medium containing 70 μg/ml ampicillin and grown at 37°C to an OD₅₅₀ of 0.1. IPTG is added to a final concentration of 2 mM and expression is induced for 1.5 h at 37°C. The cells are harvested at 4°C by centrifugation at 4000 rpm for 25 min in a Beckman GPKR centrifuge. The supermantant of CC118/pWW616 is set aside on ice for preparation of Fv(FRP5)-phoA, see Example 7.2.

7.1 Isolation of Fv(FRP5)-phoA from the periplasmic proteins of CC118/pWW616: The bacterial pellet is suspended in 10 ml TES buffer (0.2 M Tris-HCl, pH 8.0, 0.5 mM EDTA, 0.5 M sucrose) and kept on ice for 10 min. After centrifugation at 4°C for 10 min at 5000 rpm in a Heraeus minifuge, the supernatant is discarded and the washed pellet is suspended in 15 ml ice-cold TES, diluted (1:4) with water. The cells are kept on ice for 30 min and recentrifuged as above. The supernatant containing periplasmic proteins is recentrifuged at 45,000 x g for 15 min in a Beckman TL100 ultracentrifuge. The periplasmic extract is concentrated in an Amerisham ultrafiltration unit through a YM10 membrane to a final volume of 2 ml. Following fivefold dilutions with PBS and reconcentration through the YM10 membrane five times, the 1:4 diluted TES buffer of the periplasmic extract is exchanged with PBS. NaN₃ and protease inhibitors are added to the periplasmic proteins (2 ml in PBS) to the final concentration of 0.02 % NaN₃, 0.1 mM PMSF, 2 μg/ml aprotinin, 1 μg/ml leupeptin, and 1 μg/ml pepstatin. The periplasmic extract is stored at 4°C.

7.2 <u>Isolation of Fv(FRP5)-phoA from the concentrated supernatant of E. coli CC118/pWW616 cultures</u>: The supernatant (500ml) of the induced <u>E. coli</u> culture CC118/pWW616 is filtered through a 0.45 μm membrane. The filtrate is concentrated in an Amicon ultrafiltration unit through a YM10 membrane to a final volume of 10 ml in PBS as described above. NaN₃ and protease inhibitors are added to the concentrated supernatant to the final concentrations indicated above. The concentration of Fv(FRP5)-phoA in the extracts is determined by densitometry in comparison to BSA standards of coomassie stained 9 % SDS-PAGE gels.

Example 8. Activity of Fv(FRP5)-phoA

- 8.1 <u>Detection of c-erbB-2 in SKBR3 breast tumor cells by immunostaining using Fv(FRP5)-phoA</u>: The Fv domain of Fv(FRP5)-phoA enables the molecule to bind to the extracellular domain of the c-erbB-2 protein. Bound Fv(FRP5)-phoA can be visualized by staining procedures using color substrates for the detection of alkaline phosphatase activity.
- 8.1.1 <u>Fixation of cells</u>: SKBR3 human breast tumor cells carrying about 1 x 10⁶ c-erbB-2 receptors per cell are grown on fibronectin coated glass cover slips. The cells are washed twice with PBS and then fixed with PBS / 3.7 % formaldehyde at room temperature for 30 min. The fixed cells are washed three times with PBS at room temperature. Unspecific binding sites are blocked by incubating the cells for 1 h with PBS / 3 % BSA at 37°C in a humid incubator. The cells are then washed twice with PBS.
- 8.1.2 <u>Pretreatment of Fv(FRP5)-phoA:</u> Alkaline phosphatase phoA from E. <u>coli</u> must be dimerized to be enzymatically active. In the periplasm of <u>E. coli</u> natural phoA is dimerized, i.e. two molecules of phoA are held together by two Zn²⁺ ions. The Fv(FRPS)-phoA is also produced as a dimer in <u>E. coli</u>. To increase binding of Fv(FRPS)-phoA to the antigen, the dimers are monomerized by adding EGTA to the solution. This step removes Zn²⁺ from the solution. Monomerized phosphatase can be re-dimerized by the addition of Zn²⁺. EGTA is added to a final concentration of 5 mM to 200 µl of 40 x concentrated supermatant or periplasmic proteins from CC118/pWW616 (see above). The solution is incubated at 37°C for 1 h just before use in the immunoassay.
- 8.1.3 <u>Staining of cells:</u> After blocking with PBS / 3 % BSA (see above) fixed cells are incubated for 1 h with pretreated Fv(FRPS)-phoA at a concentration of 1 μg/ml at 37°C in a humidified incubator. The cells are washed three times with PBS at room temperature. The staining solution consists of 300 μl naphtol AS-MXTM phosphate (Sigma, 13 mg/ml in dimethyl formamide), 8 mg of levamisole (Sigma), and 10 mg of Fast Red TRTM salt (Sigma) added to 9.7 ml of 100 mM Tris-HCl, pH 8.2, 1 mM ZnCl₂. This mixture is prepared and filtered through a 0.45 μm filter immediately before use. ZnCl₂ is added to the staining solution to allow re-dimerization of bound Fv(FRPS)-phoA and thereby activating

the alkaline phosphatase. Cells are incubated in the Fast Red™ staining solution for 15 min at room temperature. The phosphatase activity is blocked after staining by washing the cells twice with PBS and once with 1 M KH₂PO₄. Glass cover slips are mounted with gel mount (Biomeda). The cells are examined under a fluorescence microscope using green light for excitation. Stained SKBR3 cells show intense red cell surface fluorescence.

8.2 <u>Detection of c-erbB-2 protein over-expression in immunoblots using Fv(FRP5)-phoA:</u> Proteins from total cell lysates of SKBR3 cells over-expressing c-erbB-2 protein are separated by SDS-PAGE and blotted onto PVDF membrane (Millipore). For preparation of extracts and immunoblotting technique see Example 1.3.2. Free binding sites of the membrane are blocked by incubation for 1 h at room temperature in a solution containing 10 mM Tris-HCl, pH 7.5, 0.9 % NaCl, 0.05 % Tween 20™ (BioRad), and 3 % BSA. Pretreated Fv(FRPS)-phoA (see Example 7.2.) is diluted in blocking solution to a final concentration of 0.1 µg/ml. The membrane is incubated in the Fv(FRPS)-phoA solution for 1 h at room temperature and then washed three times for 5 min at room temperature in 10 mM Tris-HCl, pH 7.5, 0.9 % NaCl, 0.05 % Tween 20™ and once in 10 mM Tris-HCl, pH 7.5, 0.9 % NaCl. For detection of bound Fv(FRPS)-phoA the membrane is incubated for 20 min at 37°C in the Fast Red™ substrate solution described in Example 7.3 without levamisole. The reaction is stopped by washing the membrane twice in water. Fv(FRP5)-phoA specifically detects the 185 kD c-erbB-2 protein.

Example 9. Expression and isolation of Fv(FRP5)-phoA from E. coli

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9.1 <u>Preparation of periplasmic extract</u>: Plasmid pWW616 is transformed into the phoA negative <u>E</u>. <u>coli</u> strain CC118 according to standard procedures (Maniatis et al., Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory, 1982). A single colony is picked and grown overnight in LB medium containing 70 µg/ml ampicillin. The overnight culture is diluted 1:10 in fresh LB medium containing ampicillin and grown at 37°C to an OD₅₆₀ of 0.1. At this point expression of the Fv(FRP5)-phoA gene is induced by the addition of IPTG to a final concentration of 2 mM, and the cells are grown for an additional 1.5 to 2 h. The cells are harvested by centrifugation and treated with a mild osmotic shock which releases the periplasmatic proteins into the supernatant. The proteins are concentrated in an Amersham ultrafiltration unit through a YM10 membrane.

9.2 Preparation of an antigen affinity column: The c-erbB-2 protein is isolated from insect cells infected with a baculovirus vector expressing the c-erbB-2 extracellular domain by standard methods (V.A. Luckow & M.D. Summers, Biotechnology 6: 47-55, 1988). MAb FSP77 is coupled to CNBR-activated Sepharose 4B™ (Pharmacia) following the instructions of the manufacturer. The insect cell lysates are incubated with the coupled MAb FSP77 in a buffer containing 50 mM Tris-HCl, pH 7.5,5 mM EGTA, 0.5 % Triton X-100™, 150 mM NaCl for 2 h at 4°C on a shaking platform. The beads are packed into a column and washed with pre-elution bufter consisting of 10 mM phosphate, pH 6.8, and 100 mM NaCl to remove non-specifically bound proteins. The c-erbB-2 protein is recovered from the column by treatment with a low pH elution buffer containing 100 mM glycine, pH 3.0, and 100 mM NaCl. The fractions from the column are collected into phosphate buffer, pH 8.0, in order to raise the pH. The c-erbB-2 extracellular domain is detected by running a part of each fraction on 8 % SDS-PAGE gel, blotting onto PVDF membrane (Millipore) and treating the filter with MAb FSP77 followed by sheep anti-mouse IgG. Bound IgG is detected by ¹²⁵I-Protein-A treatment. The fractions containing the extracellular domain are pooled and the protein is coupled to CNBR-activated Sepharose 4BTM (Pharmacia) following the instructions of the manufacturer.

9.3 <u>Isolation of Fv(FRP5)-phoA by affinity chromatography</u>: The sepharose coupled to c-erbB-2 protein (Examle 9.2) is incubated for 2-4 h at 4°C on a rocking platform with the periplasmic extract isolated as described in Example 9.1. The beads are packed into a column and washed with pre-elution buffer as in Example 9.2. The Fv(FRP5)-phoA protein is recovered by elution with the low pH elution buffer of Example 9.2. The fractions are monitored for the presence of the Fv(FRP5)-phoA by testing for phoA enzymatic activity using a standard protocol.

Example 10. Immunoassay for c-erbB-2 protein in tumors

10.1 <u>Preparation of tumor sections</u>: To determine the level of c-erbB-2 protein in tumors, tumor tissue is pretreated to give either frozen tumor sections or paraffin-embedded tumor sections. Tumor pieces are quick frozen, then cut with a cryostat, collected onto 1 % gelatin-coated glass slides, and fixed with 4 % paraformaldehyde. Following several washes with PBS, the tumor tissue sections are ready for staining. Alternatively, tumor pieces are placed in 4 % paraformaldehyde for fixation, embedded in paraffin, then sections cut and collected onto polylysine-coated glass cover slips. To prepare the sections for staining, they are heated overnight at 56°C, dewaxed in xylene, stepwise rehydrated by washing in 95 %, 70 % and 35 % ethanol and water, and washed in PBS.

10.2 <u>Pretreatment of Fv(FRP5)-phoA</u>: Since the dimer of the Fv(FRP5)-phoA as obtained from the <u>E. coli</u> periplasm does not bind optimally to the c-erbB-2 antigen, it is first monomerized. This is accomplished by treating the solution of Fv(FRP5)-phoA for 1 h at 37°C with EGTA at a final concentration of 5 mM. This treatment chelates the Zn²⁺ ions which are important for maintaining the dimeric structure of Fv(FRP5)-phoA.

10.3 Staining of the tumor sections: Non-specific staining of the tumor sections prepared according to Example 10.1 is blocked by incubating the sections in PBS containing 3 % BSA. The blocked sections are incubated for 1 - 2 h with pretreated Fv(FRP5)-phoA (Example 10.2) at a concentration of 1 μg/ml in a humidified chamber at room temperature. The sections are washed three times with PBS at room temperature. The bound Fv(FRP5)-phoA protein is detected using Fast Red™ as a substrate for the alkaline phosphatase. The staining solution consists of 300 μl naphthol AS-MX phosphate (Sigma, 13 mg/ml in dimethylformamide), 8 mg of levamisole (an inhibitor of endogenous alkaline phosphatase, Sigma), and 10 mg of Fast Red TR™ salt (Sigma) added to 9.7 ml of 100 mM Tris-HCl, pH 8.2, and 1 mM ZnCl₂. This mixture is prepared and filtered through a 0.45 μm filter immediately before use. ZnCl₂ is added to the staining solution to allow re-dimerization of the bound Fv(FRP5)-phoA protein and activation of the alkaline phosphatase. The tumor sections treated with Fv(FRP5)-phoA are incubated in the Fast Red™ staining solution for 15 min at room temperature. After staining the phosphatase activity is blocked by washing the cells twice with PBS and once with 1 M KH₂PO₄. The glass cover slips are mounted with gel mount. The cells are examined under a fluorescence microscope using green light for excitation. Positively stained cells show an intense red cell surface fluorescence.

Alternatively, the tumor sections treated with the Fv(FRP5)-phoA protein may be stained with naphthol AS-BI phosphate (Sigma) and New Fuchsin™ (Sigma), or with 5-bromo-4-chloro-3-indolyl phosphate (BCIP, Sigma) and Nitro Blue Tetrazolium™ (Sigma). The stained sections can then be viewd with a regular light microscope.

Example 11. Cloning of functional heavy and light chain rearrangements from the FWP51 hybridoma cell line

Poly(A)-containing RNA isolated from FWP51 hybridoma cells as described in Example 3.3 provides the source for cDNA synthesis and subsequent amplification of V-region minigenes, cDNA synthesis and amplification of FWP51 heavy and light chain variable domain cDNA by polymerase chain reaction is carried out as described in Example 4. Amplification products of the expected size are purified from agarose gels and cloned into appropriate vectors. Functional rearrangements are identified by sequencing.

11.1 <u>Subcloning of FWP51 heavy and light chain variable domain cDNA</u>: Material amplified according to Example 4.3 is extracted with CHCl₃ and precipitated in the presence of 200 mM LiCl. To facilitate cloning, the FWP51 heavy chain variable domain cDNA is cleaved with restriction enzymes Pstl and BstEll, the fragment purified by agarose gel electrophoresis, and ligated to Pstl and BstEll digested pWW15 DNA. The FWP51 light chain variable domain cDNA is cleaved with restriction enzymes Pvull and Bglll, the fragment is purified by agarose gel electrophoresis, and ligated to Pvull and Bglll digested pWW15 DNA (cf. Example 5). Ligation, transformation, and screening for the desired ligation products are carried out as described in Example 4.5. The following plasmids are obtained:

PCR product	Plasmid clones
Н	pWW15-VHS5-1 pWW15-VH51-2
	pWW15-VH51-3
L	pWW15-VL51-1
	pWW15-VL51-2
	pWW15-VL51-3

11.2 Sequencing: Sequencing is done as described in Example 4.6.

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Plasmids pWW15-VH5-1 (SEQ ID NO:6), pWW15-VH51-2, pWW15-VH51-3 contain identical functional SWP51 heavy chain variable domain inserts. Plasmids pWW15-VL5-1 (SEQ ID NO:7), pWW15-VL51-2, pWW15-VL51-3 contain identical functional SWP51 kappa light chain variable domain inserts. Plasmids pWW15-VH51-1 and pWW15-VL51-1 are used as a source for further subcloning steps.

Example 12. Construction of the MAb FWP51 single chain gene

12.1 <u>Assembly of the Fv fusion gene:</u> Plasmid pWW15-VH51-1 is digested with PstI and BstEII and the 342 bp heavy chain variable domain fragment of FWP51 is isolated. It is cloned into PstI/BstEII digested pWW15-VL-51-1 yielding the plasmid pWW15-Fv51 (SEQ ID NO:8).

12.2 <u>Mutation of the single-chain Fv(FWP51) gene:</u> To allow gene fusion between the single-chain Fv(SWP51) encoding gene from pWW15-Fv51 and effector genes the stop codon at sequence position 729 to 731 in pWWFv15-51 (SEQ ID NO:8) is deleted as follows (see also Example 6.1): plasmid DNA of pWW15-Fv51 is digested with BstEll and

Bglll and the linker sequence and SWP51 light chain variable domain encoding fragment is isolated. In another digestion, pWW15-Fv51 is cleaved with BstEll and Bcll. Thus, the large fragment containing vector sequences and the FWP51 heavy chain variable domain encoding sequence is isolated. The BstEll/Bglll V_L fragment is now inserted into BstEll/Bcll cleaved pWW15-Fv51 containing V_H. The resulting plasmid pWW15-Fv51-ORF is used as a source for th construction of Fv(FWP51)-effector fusion genes.

Example 13. Construction of single-chain Fv-exotoxin A fusion gene expression plasmids

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The MAb FRPS and MAb FWP51 single-chain Fv genes are fused to a truncated bacterial toxin, exotoxin A (ETA) from <u>Pseudomonas aeruginosa</u>. These chimeric genes encode recombinant immunotoxins which selectively inhibit protein synthesis in c-erbB-2 expressing cells.

13.1 <u>Mutation of the Exotoxin A gene of Pseudomonas aeruginosa PAK:</u> For the construction of Fv-exotoxin A (Fv-ETA) fusion genes the ETA gene from <u>Pseudomonas aeruginosa</u> PAK is mutated to delete the original cell binding domain I at the N-terminus of the toxin and to generate a Xbal cleavage site at the former domain I/domain II boundary of the ETA coding region. Plasmid pMS150A (Lory et al., J. Bacteriol. 170: 714, 1988) is linearized by EcoRI cleavage. 20 ng of the linearized template DNA is used for a 100 μI PCR reaction carried out as described previously using the following oligonucleotides as primers 1 and 2.

1: 5'-CACGGAAGCTTAAGGAGATCTGCATGCTTCTAGAGGGCGGCA-

GCCTGGCCGCGCTG-3'

2: 5'-GCGGATCGCTTCGCCCAGGT-3'

Following HindIII/Sall digestion of the PCR products, a 201 bp fragment is isolated from a 1.5% agarose gel and inserted into HindIII/Sall digested plasmid pUC18. Ligation is carried out as described above. Ligated DNA is transformed into E.coli XL1 Blue™ (Stratagene). Two recombinant plasmids are isolated and the insert DNA is sequenced as described above using pUC universal and reverse primers (Boehringer). One plasmid containing the expected product is designated pWW22 (SEQ ID NO:9) and used as a source for further subcloning steps. Plasmid pWW22 is cleaved with HindIII and Sall, the mutated ETA gene fragment is isolated, and inserted into the large fragment of HindIII/ Sall digested plasmid pMS150A containing pUC9 vector sequences and part of the ETA gene coding for the C-terminal half of the toxin. Thereby in the resulting plasmid pWW20 a truncated ETA gene coding for domains II and III of the toxin is created.

13.2 <u>Assembly of single-chain Fv-ETA lusion genes</u>: HindllI/Xbal single-chain Fv gene fragments suitable for the construction of Fv-ETA lusion genes are isolated from plasmid pWW53 (single-chain Fv FRP5), and plasmid pWW15-Fv51-ORF (single-chain Fv FWP51) and inserted into HindllI/Xbal digested pWW20. Ligation and transformation into E.coli XL1 Blue™ (Stratagene) are carried out as described above. The resulting plasmids pWW20-Fv5 (Fv(FRP5)-ETA) and pWW20-Fv51 (Fv(FWP51)-ETA) are used as a source for further subcloning steps.

13.3 Construction of single-chain Fv-exotoxin A fusion gene expression plasmids: For the expression of single-chain Fv-exotoxin A fusion genes in E.coli the expression plasmid pFLAG-1 (IBI Biochemicals) is used. The fusion-genes are fused in frame to the outer membrane protein A (ompA) signal sequence encoded by pFLAG-1. Plasmid DNA from pWW20-Fv5 and pWW20-Fv51 is digested with HindIII and blunt ends are created by Klenow fill-in as described in Example 4.5. Blunt ended DNA is digested with EcoRI and single-chain Fv-ETA gene fragments are isolated (Fv(FRP5)-ETA: 1916 bp, Fv(FWP51)-ETA: 1916 bp). pFLAG-1 plasmid DNA is digested with HindIII, blunt ends are created as described above, the resulting DNA fragment is isolated, and digested with EcoRI. Blunt-end/ EcoRI Fv-ETA fusion gene fragments are inserted into the modified pFLAG-1 plasmid DNA. Thereby Fv-ETA fragments are fused in frame to the ompA signal sequence of pFLAG-1 creating plasmids pWW215-5 for the expression of Fv (FRP5)-ETA (SEQ ID NO: 10) and pWW215-51 for the expression of Fv(FWP51)-ETA (SEQ ID NO: 11).

Example 14. Expression and isolation of Fv(FRP5)-ETA and Fv(FWP51)-ETA from E. coli

14.1 <u>Preparation of total lysates</u>: Plasmids pWW215-5 and pWW215-51 are transformed into the <u>E.coli</u> strain CC118 according to standard procedures (see Example 9.1). Single colonies are picked and grown ovemight in LB medium containing 100 μg/ml ampicillin and 0.4% glucose. The overnight cultures are diluted 1:30 in fresh LB medium containing ampicillin and glucose and grown at 37°C to an OD₅₅₀ of 0.5. At this point expression of the Fv(FRP5)-ETA and Fv(FWP51)-ETA genes is induced by the addition of IPTG to a final concentration of 0.5 mM, and the cells are grown for an additional 30 min. The cells are harvested by centrifugation and lysed by sonication in PBS/1 mM CaCl₂. The lysates are cleared by ultracentrifugation at 25 000 g for 45 min at 4°C. The supernatants are collected.

14.2 Isolation of Fv(FRP5)-ETA and Fv(FWP51)-ETA by affinity chromatography: Cleared E.coli lysates containing

the 66.4 kDa Fv(FRP5)-ETA or the 66.3 kDa Fv(FWP51)-ETA protein are passed through a M1 monoclonal antibody affinity column (IBI Biochemicals). The column is washed three times with PBS/1 mM CaCl₂. Bound Fv(FRP5)-ETA or Fv(FWP51)-ETA proteins are eluted with PBS/2 mM EDTA. The fractions are monitored for the presence of Fv-ETA proteins by SDS-PAGE and immunoblotting (see Example 1.3.2) using an anti-exotoxin A antiserum developed in rabbit

Example 15. <u>Selective inhibition of protein synthesis in c-erbB-2 expressing cells with Fv(FRP5)-ETA and Fv(FWP51)-ETA</u>

In vitro the recombinant immunotoxins Fv(FRP5)-ETA and Fv(FWP51)-ETA selectively inhibit protein synthesis and growth of cells expressing high levels of the human c-erbB-2 protein. The immunotoxins do not affect cells expressing no, or low levels of human c-erbB-2 protein.

15.1 Immunotoxin treatment of cell lines: Human breast and ovarian tumor cell lines SK-BR3, MDAMB-231, MDA-MB-453, HTB77, the mouse mammary epithelial cell line HC11, and HC11 cells transfected with the human c-erbB-2 cDNA are plated on 48 well tissue culture plates (Costar) at a density of 105⁵ cells/well. After 4 h the medium is removed and replaced by normal growth medium containing Fv(FRP5)-ETA or Fv(FWP51)-ETA at various concentrations ranging from 1 to 1000 ng/ ml. The cells are incubated with toxin fusion proteins for 16 h.

15.2 3H-leucine labeling of cells: The immunotoxin-treated cells are washed twice and incubated in normal growth medium containing 4 µCi 3H-leucin/ ml for 4 h. The labeled cells are washed twice and 3H-leucine labeled total proteins are harvested by TCA precipitation onto Whatman GFC filters. The rate of protein synthesis in immunotoxin-treated cells is determined in comparison to untreated control cells.

Example 16: Fv(FRP5)-ETA and MAbs FWP51 and FSP77 inhibit the growth of c-erbB-2 expressing cells in nude mice.

The administration of Fv(FRP5)-ETA and the MAbs FWP51 and FSP77 to animals injected with c-erbB-2 expressing cells inhibits the tumor growth of these cells.

16.1 Nude mouse tumor model.: The NIH/3T3 mouse fibroblast cell line is transfected according to conventional, previously described methods (Graham & van der Eb, Virology 52: 456, 1973) with a plasmid expressing the point mutated, activated human c-erbB-2 protein (Masuko et al., Jpn. Cancer Res. 80: 10, 1989) and with the plasmid pSV2neo (Southem & Berg, J. Mol. Appl. Genet. 1:327, 1982) which encodes the gene for resistance to the drug G418. Transfected cells are selected 2 weeks in medium containing 500 ug/ml G418 (Geneticin, Gibco-BRL). Individual clones are selected and analyzed for the expression of the human c-erbB-2 protein using conventional protein blotting techniques (Towbin et al., Proc. Natl. Acad. Sci. USA 76: 4350, 1979). A clone expressing moderate levels of the point mutated, activated human c-erbB-2 protein (clone 3.7) is selected, and tested for growth in nude mice. 2-5 x 106 clone 3.7 cells (per animal) suspended in 0.2 ml PBS are subcutaneously injected into the flank of female Balb/c nude mice. The 3.7 cells injected at a dose of 2 x 106 cells rapidly form tumors in nude mice (control animals, cf. Example 16.2)

16.2 <u>Immunotoxin treatment of animals</u>: 2 x 106 clone 3.7 cells are injected subcutaneously into nude mice. The animals are treated continuously for a total of 7 days with the Fv(FRP5)-ETA. 200 μl of Fv(FRP5)-ETA (concentration 35 μg/ml in PBS) is placed in an osmotic pump (Alzet mini osmotic pump, Model 2001, Alza, Palo Alto, CA, #94303-0802) which is implanted subcutaneously into the animals at the same time as the clone 3.7 cells are injected. The pump continuously releases Fv(FRP5)-ETA and delivers 1 μg/day for 7 days to each animal. In comparison with the control animals (cf. Example 16.1), the administration of Fv(FRP5)-ETA delays the onset of tumor formation.

16.3 MAb treatment of animals: 5 x 106 clone 3.7 cells are injected subcutaneously into nude mice. Starting on the same day as injection of clone 3.7 cells, the animals are treated daily, for a total of 10 days, with either MAb FWP51 or MAb FSP77 (MAb dose is 50 ug/200 ul BSS/day). The MAb is injected intraveneously in the tail vein of the mouse. Both antibodies delay the onset of tumor growth. Compared therewith, a synergistic effect in inhibiting tumor growth is observed on simultaneous administration of both antibodies MAb FWP51 and MAb FSP77.

Sequence listing

SEQ ID NO: 1

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SEQUENCE TYPE: nucleotide
SEQUENCE LENGTH: 361 bp
MOLECULE TYPE: plasmid DNA
ORIGINAL SOURCE ORGANISM: mouse
IMMEDIATE EXPERIMENTAL SOURCE: <u>E</u>. <u>coli</u>
NAME OF CELL CLONE: pMZ16/1

FEATURES:	from 6 to 27 bp	VH1BACK primer region
	from 95 to 109 bp	CDR _{1H}
	from 152 to 202 bp	CDR _{2H}
	from 299 to 328 bp	CDR _{3H}
	from 329 to 361 bp	VH1FOR primer region

PROPERTIES: encodes the heavy chain variable domain of monoclonal antibody FRPS

TCTAGAGGTG	AAACTGCAGC	AGTCTGGACC	TGAACTGAAG	AAGCCTGGAG	50
AGACAGTCAA	GATCTCCTGC	AAGGCCTCTG	GGTATCCTTT	CACAAACTAT	100
GGAATGAACT	GGGTGAAGCA	GGCTCCAGGA	CAGGGTTTAA	AGTGGATGGG	150
CTGGATTAAC	ACCTCCACTG	GAGAGTCAAC	ATTTGCTGAT	GACTTCAAGG	200
GACGGTTTGA	CTTCTCTTTG	GAAACCTCTG	CCAACACTGC	CTATTTGCAG	250
ATCAACAACC	TCAAAAGTGA	AGACATGGCT	ACATATTTCT	GTGCAAGATG	300
GGAGGTTTAC	CACGGCTACG	TTCCTTACTG	GGGCCAAGGG	ACCACGGTCA	350
CCGTCTCCTC	A				361

SEQ ID NO:2

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SEQUENCE TYPE: nucleotide SEQUENCE LENGTH: 407 bp MOLECULE TYPE: plasmid DNA ORIGINAL SOURCE ORGANISM: mouse

IMMEDIATE EXPERIMENTAL SOURCE: E. coli

NAME OF CELL CLONE: pMZ18/1

FEATURES:	from 6 to 28 bp	MCK2 primer region
	from 98 to 130 bp	CDR _{1L}
	from 176 to 196 bp	CDR _{2L}
	from 293 to 319 bp	CDR _{3L}
	from 374 to 404 bp	MCK2 primer region

PROPERTIES: encodes the kappa light chain variable domain of monoclonal antibody FRPS

TCTAGTCACT GGATGGTGGG AAGATGGAGA CATTGTGATG ACCCAGTCTC	50
ACAAATTCCT GTCCACTTCA GTAGGAGACA GGGTCAGCAT CACCTGCAAG	100
GCCAGTCAGG ATGTGTATAA TGCTGTTGCC TGGTATCAAC AGAAACCAGG	150
ACAATCTCCT AAACTTCTGA TTTACTCGGC ATCCTCCCGG TACACTGGAC	200
TCCCTTCTCG CTTCACTGGC AGTGGCTCTG GGCCGGATTT CACTTTCACC	250
ATCAGCAGTG TGCAGGCTGA AGACCTGGCA GTTTATTTCT GTCAGCAACA	300
TTTTCGTACT CCATTCACGT TCGGCTCGGG GACAAAATTG GAAATAAAA	350
GGGCTGATGC TGCACCAACT GTATCCATCT TCCCACCATC CAGTGACTAC	3 400
AACTAGA	407

SEQ ID NO:3

SEQUENCE TYPE: nucleotide SEQUENCE LENGTH: 175 bp

MOLECULE TYPE: plasmid DNA
ORIGINAL SOURCE ORGANISM: fully synthetic IMMEDIATE EXPERIMENTAL SOURCE: <u>E. coli</u> NAME OF CELL CLONE: pWW19

FEATURES:	from 30 to 35 bp	Pstl site
	from 38 to 44 bp	BstEll site for subcloning of heavy chain variable domain
	from 54 to 98 bp	coding sequence of (GlyGlyGlyGlySer)3 linker
	from 105 to 110 bp	Pvull site
	from 112 to 117 bp	BgIII site
	from 120 to 125 bp	Bcll site for subcloning of light chain variable domain

_	AAGCTTGCAT	GCAAGCTTCT	CAGGTACAAC	TGCAGGAGGT	CACCGTTTCC	50
5	TCTGGCGGTG	GCGGTTCTGG	TGGCGGTGGC	TCCGGCGGTG	GCGGTTCTGA	100
10	CATCCAGCTG	GAGATCTAGC	TGATCAAAGC	TCTAGAGGAT	CCCCGGGTAC	150
	CGAGCTCGAA	TTCACTGGCC	GTCGT			175

SEQ ID NO:4

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SEQUENCE TYPE: nucleotide with corresponding protein SEQUENCE LENGTH: 748 bp MOLECULE TYPE: plasmid DNA ORIGINAL SOURCE ORGANISM: mouse IMMEDIATE EXPERIMENTAL SOURCE: E. coli NAME OF CELL CLONE: pWW52

FEATURES:	from 1 to 8 bp	synthetic spacer
	from 9 to 365 bp	FRP5 heavy chain variable domain
	from 99 to 113 bp	CDR _{1H}
	from 156 to 206 bp	CDR _{2H}
	from 303 to 332 bp	CDR _{3H}
	from 366 to 410 bp	15 amino acids linker sequence
	from 411 to 728 bp	FRP5 light chain variable domain
	from 480 to 512 bp	CDR _{1L}
	from 558 to 578 bp	CDR _{2L}
	from 675 to 701 bp	CDR _{at}

PROPERTIES: Fv heavy chain/light chain variable domain fusion protein binding to the extracellular domain of the growth factor receptor c-erbB-2

	AAGCT T	CT C	AG GT	ra ca	AA CI	rg CI	AG C	AG TC	T GO	SA CC	T G	AA C	ľG	41
	S	er G	ln Va	al Gl	n Le	eu Gl	ln Gl	ln Se	er Gl	ly Pr	co G	lu Le	eu	
5					5	5				1	LO			
	AAG AAG	CCT	GGA	GAG	ACA	GTC	AAG	ATC	TCC	TGC	AAG	GCC	TCT	83
10	Lys Lys	Pro	Gly	Glu	Thr	Val	Lys	Ile	Ser	Cys	Lys	Ala	Ser	
		15					20					25		
15	GGG TAT	CCT	TTC	ACA	AAC	TAT	GGA	ATG	AAC	TGG	GTG	AAG	CAG	125
,,	Gly Tyr	Pro	Phe	Thr	Asn	Tyr	Gly	Met	Asn	Trp	Val	Lys	Gln	
	_		30					35					40	
20	GCT CCA	GGA	CAG	GGT	TTA	AAG	TGG	ATG	GGC	TGG	ATT	AAC	ACT	167
	Ala Pro	Gly	Gln	Gly	Leu	Lys	Trp	Met	Gly	Trp	Ile	Asn	Thr	
				45					50					
25														

	TCC	ACT	GGA	GAG	TCA	ACA	TTT	GCT	GAT	GAC	TTC	AAG	GGA	CGG	209
	Ser	Thr	Gly	Glu	Ser	Thr	Phe	Ala	Asp	Asp	Phe	Lys	Gly	Arg	
5	55					60					65				
	TTT	GAC	TTC	TCT	TTG	GAA	ACC	TCT	GCC	AAC	ACT	GCC	TAT	TTG	251
10	Phe	Asp	Phe	Ser	Leu	Glu	Thr	Ser	Ala	Asn	Thr	Ala	Tyr	Leu	
		70					75					80			
15	CAG	ATC	AAC	AAC	CTC	AAA	AGT	GAA	GAC	ATG	GCT	ACA	TAT	TTC	293
	Gln	Ile	Asn	Asn	Leu	Lys	Ser	Glu	Asp	Met	Ala	Thr	Tyr	Phe	
			85					90				•	95	•	
20	TGT	GCA	AGA	TGG	GAG	GTT	TAC	CAC	GGC	TAC	GTT	CCT	TAC	TGG	335
	Cys	Ala	Arg	Trp	Glu	Val	Tyr	His	Gly	Tyr	Val	Pro	Tyr	Trp	
				100					105					110	
25															
	GGC	CAA	GGG	ACC	ACG	GTC	ACC	GTT	TCC	TCT	GGC	GGT	GGC	GGT	377
	Gly	Gln	Gly	Thr	Thr	Val	Thr	Val	Ser	Ser	Gly	Gly	Gly	Gly	
30					115					120					
	TCT	GGT	GGC	GGT	GGC	TCC	GGC	GGT	GGC	GGT	TCT	GAC	ATC	CAG	419
35	Ser	Gly	Gly	Gly	Gly	Ser	Gly	Gly	Gly	Gly	Ser	Asp	Ile	Gln	
	125					130					135				
40	CTG	ACC	CAG	TCT	CAC	AAA	TTC	CTG	TCC	ACT	TCA	GTA	GGA	GAC	461
40	Leu	Thr	Gln	Ser	His	Lys	Phe	Leu	Ser	Thr	Ser	Val	Gly	Asp	
		140					145					150			
45													TAT		503
	Arg	Val	Ser	Ile	Thr	Cys	Lys		Ser	Gln	Asp	Val	Tyr	Asn	
			155			•		160					165		
50															
														AAA	545
	Ala	Val	Ala	_	Tyr	Gln	Gln	Lys			Gln	Ser	Pro	Lys	
				170					175					180	

	CTT	CTG	ATT	TAC	TCG	GCA	TCC	TCC	CGG	TAC	ACT	GGA	GTC	CCT	587
	Leu	Leu	Ile	Tyr	Ser	Ala	Ser	Ser	Arg	Tyr	Thr	Gly	Val	Pro	
5					185					190					
	TCT	CGC	TTC	ACT	GGC	AGT	GGC	TCT	GGG	CCG	GAT	TTC	ACT	TTC	629
10	Ser	Arg	Phe	Thr	Gly	Ser	Gly	Ser	Gly	Pro	Asp	Phe	Thr	Phe	
	195					200					205				
15	ACC	ATC	AGC	AGT	GTG	CAG	GCT	GAA	GAC	CTG	GCA	GTT	TAT	TTC	671
	Thr	Ile	Ser	Ser	Val	Gln	Ala	Glu	Asp	Leu	Ala	Val	Tyr	Phe	
		210					215					220		-	
20										٠					
	TGT	CAG	CAA	CAT	TTT	CGT	ACT	CCA	TTC	ACG	TTC	GGC	TCG	GGG	713
	Cys	Gln	Gln	His	Phe	Arg	Thr	Pro	Phe	Thr	Phe	Gly	Ser	Gly	
			225					230					235		
25															
	ACA	AAA	TTG	GAG	ATC	TAG	CTGAT	rca .	AAGC'	TCTA	GA				748
	Thr	Lys	Leu	Glu	Ile										
30				240											
	SEQ ID N	IO:5													
35		UENCE				h corre	spondin	g prote	in						
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PROPERTIES: Fv heavy chain/light chain variable domain and alkaline phosphatase fusion protein Fv(FRPS)-phoA binding to the growth factor receptor c-erbB-2

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5									AAA Lys -20				43
10				GCA Ala									85 _.
15				CAA Gln									127
20				ACA Thr									169
25		Phe		AAC Asn			Met				Gln		211
<i>30</i> <i>35</i>		30					35				40		
40													
45													
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	GGA	CAG	GGT	TTA	AAG	TGG	ATG	GGC	TGG	ATT	AAC	ACC	TCC	ACT	253
	Gly	Gln	Gly	Leu	Lys	Trp	Met	Gly	Trp	Ile	Asn	Thr	Ser	Thr	
5			45					50					55		
	•														
	GGA	GAG	TCA	ACA	TTT	GCT	GAT	GAC	TTC	AAG	GGA	CGG	TTT	GAC	295
10	Gly	Glu	Ser	Thr	Phe	Ala	Asp	Asp	Phe	Lys	Gly	Arg	Phe	Asp	
				60					65					70	
	TTC	TCT	TTG	GAA	ACC	TCT	GCC	AAC	ACT	GCC	TAT	TTG	CAG	ATC	337
15	Phe	Ser	Leu	Glu	Thr	Ser	Ala	Asn	Thr	Ala	Tyr	Leu	Gln	Ile	
					75					80				•	
20													TGT		379
	Asn	Asn	Leu	Lys	Ser	Glu	Asp	Met	Ala	Thr	Tyr	Phe	Cys	Ala	
	85					90					95				
25															
													GGC		421
	Arg	Trp	Glu	Val	Tyr	His	Gly	Tyr	Val	Pro	Tyr		Gly	Gln	
30		100					105					110			
													TCT		463
	Gly	Thr		Val	Thr	Val	Ser	Ser	Gly	Gly	Gly	Gly	Ser	Gly	
35			115					120					125		
													CTG		505
40	Gly	Gly	Gly		Gly	Gly	Gly	Gly		Asp	Ile	Gin	Leu		
				130					135					140	
												~~~		000	547
45													AGG		547
	Gln	Ser	His	Lys		Leu	Ser	Thr	Ser		GIY	Asp	Arg	vaı	
					145					150					
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	155					160					165				
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	GCC	TGG	TAT	CAA	CAG	AAA	CCA	GGA	CAA	TCT	CCT	AAA	CTT	CTG	631
	Ala	Trp	Tyr	Gln	Gln	Lys	Pro	Gly	Gln	Ser	Pro	Lys	Leu	Leu	
5		170					175					180			
	•														
	ATT	TAC	TCG	GCA	TCC	TCC	CGG	TAC	ACT	GGA	GTC	CCT	TCT	CGC	673
10	Ile	Tyr	Ser	Ala	Ser	Ser	Arg	Tyr	Thr	Gly	Val	Pro	Ser	Arg	
			185					190					195		
													ACC		715
15	Phe	Thr	Gly	Ser	Gly	Ser	Gly	Pro	Asp	Phe	Thr	Phe	Thr	Ile	
				200					205					21.0	
20													TGT		757
	Ser	Ser	Val	Gln	Ala	Glu	Asp	Leu	Ala		Tyr	Phe	Cys	Gln	
	•				215					220					
25															
													ACA		799
	Gln	His	Phe	Arg	Thr		Phe	Thr	Phe	Gly		Gly	Thr	Lys	
30	225					230					235				
-															
													CGG		841
	Leu		Ile	Lys	Ala	Leu		Pro	Val	Leu	Glu		Arg	Ala	
35		240					245					250			
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													CGT		883
40	Ala	Gin	_	Asp	11e	Thr	Ala		GIY	GIĀ	Ата	Arg	Arg	ren	
			255					260					265		
		000	63.M	636	<b>.</b>	666	ccm	CITIC	ccm	~ » m	mc m	COO	200	CATI	925
45													AGC Ser		943
	Thr	GIĀ	ASD	270	THE	Ата	АТА	rea	275	мър	Ser	Leu	261	280	
				270					275					200	
50	222	CCM	GC A	222	እአጥ	יוושרוי ע	יוייית ע	تكليك	CTC	עושוי ע	GGC	ርኔጥ	GGG	ATYC	967
													Gly		501
	ъÃЗ	FIO	vra	ny s	285	116	116	Leu	Jeu	290	GIY	nop			
					203					230					
55															

	GGG	GAC	TCG	GAA	TTA	ACT	GCC	GCA	CGT	AAT	TAT	GCC	GAA	GGT	1009
	Gly	Asp	Ser	Glu	Ile	Thr	Ala	Ala	Arg	Asn	Tyr	Ala	Glu	Gly	
5	295					300					305				
	GCG	GGC	GGC	TTT	TTT	AAA	GGT	ATA	GAT	GCC	TTA	CCG	CTT	ACC	1051
10	Ala	Gly	Gly	Phe	Phe	Lys	Gly	Ile	Asp	Ala	Leu	Pro	Leu	Thr	
,,,		310					315					320			
	GGG	CAA	TAC	ACT	CAC	TAT	GCG	CTG	AAT	AAA	AAA	ACC	GGC	AAA	1093
15	Gly	Gln	Tyr	Thr	His	Tyr	Ala	Leu	Asn	Lys	Lys	Thr	Gly	Lys	
			325					330					335		
20	CCG	GAC	TAC	GTC	ACC	GAC	TCG	GCT	GCA	TCA	GCA	ACC	GCC	TGG	1135
	Pro	Asp	Tyr	Val	Thr	Asp	Ser	Ala	Ala	Ser	Ala	Thr	Ala	Trp	
				340					345					350	
25															
	TCA	ACC	GGT	GTC	AAA	ACC	TAT	AAC	GGC	GCG	CTG	GGC	GTC	GAT	1177
	Ser	Thr	Gly	Val	Lys	Thr	Tyr	Asn	Gly	Ala	Leu	Gly	Val	Asp	
					355					360					
30															
	ATT	CAC	GAA	AAA	GAT	CAC	CCA	ACG	ATT	CTG	GAA	ATG	GCA	AAA	1219
	Ile	His	Glu	Lys	Asp	His	Pro	Thr	Ile	Leu	Glu	Met	Ala	Lys	
35	365					370					375				
	GCC	GCA	GGT	CTG	GCG	ACC	GGT	AAC	GTT	TCT	ACC	GCA	GAG	TTG	1261
40	Ala	Ala	Gly	Leu	Ala	Thr	Gly	Asn	Val	Ser	Thr	Ala	Glu	Leu	
		380					385					390			
45	CAG	GAT	GCC	ACG	CCC	GCT	GCG	CTG	GTG	GCA	CAT	GTG	ACC	TCG	1303
45	Gln	Asp	Ala	Thr	Pro	Ala	Ala	Leu	Val	Ala	His	Val	Thr	Ser	
			395					400					405		
50	CGC	AAA	TGC	TAC	GGI	CCG	AGC	GCG	ACC	AGT	GAA	AAA	TGT	CCG	1345
														Pro	
	- 3	• -	•	410					415			_		420	

	GGT	AAC	GCT	CTG	GAA	AAA	GGC	GGA	AAA	GGA	TCG	ATT	ACC	GAA	1387
	Gly	Asn	Ala	Leu	Glu	Lys	Gly	Gly	Lys	Gly	Ser	Ile	Thr	Glu	
5					425					430					
	CAG	CTG	CTT	AAC	GCT	CGT	GCC	GAC	GTT	ACG	CTT	GGC	GGC	GGC	1429
	Gln	Leu	Leu	Asn	Ala	Arg	Ala	Asp	Val	Thr	Leu	Gly	Gly	Gly	
10	435					440					445				
	GCA	AAA	ACC	TTT	GCT	GAA	ACG	GCA	ACC	GCT	GGT	GAA	TGG	CAG	1471
15			Thr												
		450					455				_	460	_		
20	GGA	AAA	ACG	CTG	CGT	GAA	CAG	GCA	CAG	GCG	CGT	GGT	ТАТ	CAG	1513
			Thr												
	013	-1 -	465		3	•		470					475		
			.03												
25	വപറ	GTG	AGC	GAT	ССТ	GCC	ጥሮል	CTG	ልልጥ	TCG	GTG	ACG	GAA	GCG	1555
			Ser												
	БСС	V41	JCI	480	1114	****	501	200	485	001			014	490	
30				400					103					-50	
	2.20	CAC	CAA	***	ccc	CTC	CTT	ccc	CTC	بالجلجل	CCT	GAC	ccc	ААТ	1597
			Gln												133.
35	ASII	GIII	GIII	гуs	495	Deu	Deu	Gry	neu	500	AIG	дон	GIY	YOU	
55					493					200					
	3.000	223	OMO	000	maa	CMN	CCA	ccc	222	CCA	»CC	ma.c	CAM	GGC	1639
															1033
40		Pro	Val	Arg	TIP		GIY	PIO	гуу	AIG		IAT	птэ	GIA	
	505					510					515				
						~~`	C.T.C.				221		666	~~~	1.601
45			GAT												1681
	Asn		Asp	Lys	Pro	Ala		inr	Cys	ınr	Pro		PIO	GIN	
		520					525					530			
50															1500
50														AAA	1723
	Arg	Asn	Asp	Ser	Val	Pro	Thr		Ala	Gln	Met	Thr		Lys	
-			535					540					545		
55															

......

	GCC	ATT	GAA	TTG	TTG	AGT	AAA	AAT	GAG	AAA	GGC	TTT	TTC	CTG	1765
	Ala	Ile	Glu	Leu	Leu	Ser	Lys	Asn	Glu	Lys	Gly	Phe	Phe	Leu	
5				550					555					560	
	CAA	GTT	GAA	GGT	GCG	TCA	ATC	GAT	AAA	CAG	GAT	CAT	GCT	GCG	1807
	Gln	Val	Glu	Gly	Ala	Ser	Ile	Asp	Lys	Gln	Asp	His	Ala	Ala	
10					565					570				•	
	ААТ	ССТ	TGT	GGG	CAA	ATT	GGC	GAG	ACG	GTC	GAT	CTC	GAT	GAA	1849
15					Gln										
	575		٠,٠	,	•	580	•				585		_		
	3,3														
20	GCC	СТА	CAA	CGG	GCG	CTG	GAA	TTC	GCT	AAA	AAG	GAG	GGT	AAC	1891
					Ala										
	niu	590	01	9			595					600	-		
		550					555								
25	»CC	CIPC	CTC	ልጥል	GTC	ACC	CCT	САТ	CAC	GCC	CAC	GCC	AGC	CAG	1933
					Val										
	1111	Dea	605		Vai	1111	niu	610				•••	615		
30			003					010					010		
	» mm	Cmm	CCC	ccc	GAT	»CC	222	ርርጥ	CCG	GGC	CTC	ACC	CAG	GCG	1975
					Asp										
35	116	Val	MIG	620	ASP	1111	Ly S	AIG	625	GIY	·	1112	<u> </u>	630	
55				020					023					050	
	C/M3	3 3 00	300	222	CAM	ccc	CCA	CTC	איזייני	GTG	ልጥር	ልርጥ	ጥልሮ	GGG	2017
					Asp										2017
40	Leu	ASII	THE	пÃ2		GIŞ	AIG	VOI	Mec	640	Mec	Jer	171	O13	
					635					040					
		mcc.	CAA	CAC	CNE	ייים א	CAA	CAA	CAT	እርር	ccc	እርጥ	CAG	TTG	2059
45														Leu	2033
			GIU	GIU	Asp		GIII	Giu	птр	1111		Ser	GIII	neu	
	645					650					655				
50				000	m	000	000	C1 C	000	000	220	Citture	COM	CCN	2101
50														GGA	2101
	Arg			Ala	Tyr	GIY			Ala	ATA	ASN		val	GIÀ	
		660					665					670			

5		Thr Asp Leu	TTC TAC ACC ATG A Phe Tyr Thr Met L 680		2143
10	CTG GGG CTG AAA Leu Gly Leu Lys	TAAAACCGCG C	CCGGCAGTG AATTTTC	GCT	2185
	690 GCCGGGTGGT TTTT	TTGCTG TTAGCA	ACCA GACTTAATGG C	AGAGCTC	2233
15	SEQ ID NO:6				
20	SEQUENCE TYPE: nuclei SEQUENCE LENGTH: 34 MOLECULE TYPE: plasm ORIGINAL SOURCE ORG IMMEDIATE EXPERIMEN NAME OF CELL CLONE:	2 bp id DNA iANISM: mouse TAL SOURCE: E.∞li			
25	FEATURES:	from 1 to 14 bp from 82 to 96 bp from 139 to 189 bp	partial sequence of VH1BAC CDR _{1H} CDR _{2H}	K primer region	
30		from 286 to 318 bp from 317 to 342 bp	CDR _{3H} partial sequence of VH1FOF	primer region	
35		-	domain of monoclonal antibo	-	50
	GTCCTGCAAG GC	TTCTGATT ACAC	CCTTCAC CAGCTACTG	G ATGAACTGGG	100
40	TGAAGCAGAG GC	CTGGACAA GGC	CTTGAAT GGATTGGTA	r gattgatcci	150
	TCAGACAGTG AA	ACTCAATA CAA	CAAATG TTCAAGGAC	A AGGCCGCATT	200
45	GACTGTAGAC AA	GTCCTCCA ATA	CAGCCTA CATGCAACT	C AGCAGCCTGA	A 250
50	CATCTGAGGA CI	CTGCGGTC TAT	PACTGTG CAAAAGGGG	G GGCCTCTGGG	300
	GACTGGTACT TO	CGATGTCTG GGGG	CCAAGGG ACCACGGTC	A CC	342
55	SEQ ID NO:7				

39

SEQUENCE TYPE: nucleotide

SEQUENCE LENGTH: 310 bp
MOLECULE TYPE: plasmid DNA
ORIGINAL SOURCE ORGANISM: mouse
IMMEDIATE EXPERIMENTAL SOURCE: E.coli

NAME OF CELL CLONE: pWW15-VL51-1

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FEATURES:	from 1 to 18 bp	partial sequence of VK1BACK primer region
	from 64 to 96 bp	CDRiL
	from 142 to 162 bp	CDR _{2L}
	from 259 to 282 bp	CDR _{3L}
	from 292 to 310 bp	partial sequence of VK1FOR primer region

PROPERTIES: encodes the light chain variable domain of monoclonal antibody FWP51

15	Properties. encodes the light chain variable contain of monocolar animody PWF51	
13	CAGCTGACCC AGTCTCCATC CTCACTGTCT GCATCTCTGG GAGGCGAAGT	50
20	CACCATCACT TGCAAGGCAA GCCAAGACAT TAAGAAGTAT ATAGCTTGGT	100
	ACCAACACA GCCTGGAAAA AGTCCTCGGC TACTCATACA CTACACATCT	150
25	GTATTACAGC CAGGCATCCC ATCCAGGTTC AGTGGAAGTG GGTCTGGGAG	200
30	AGATTATTCC TTCAGCATCC ACAACCTGGA GCCTGAAGAT ATTGCAACTT	250
	ATTATTGTCT ACATTATGAT TATCTGTACA CGTTCGGAGG GGGCACCAAG	300
35	CTGGAGATCT	310

## SEQ ID NO:8

50

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40 SEQUENCE TYPE: nucleotide with corresponding protein SEQUENCE LENGTH: 748 bp
MOLECULE TYPE: plasmid DNA ORIGINAL SOURCE ORGANISM: mouse
IMMEDIATE EXPERIMENTAL SOURCE: E.coli
NAME OF CELL CLONE: pWW15-Fv51

FEATURES:	from 1 to 8 bp	synthetic spacer
	from 9 to 368 bp	FWP 51 heavy chain variable domain
	from 99 to 113 bp	CDR1H
	from 156 to 206 bp	CDR2H
	from 303 to 335 bp	CDR3H
	from 369 to 413 bp	synthetic spacer
	from 414 to 728 bp	FWP 51 light chain variable domain
	from 483 to 515 bp	CDR1L
	from 561 to 581 bp	R2L
	from 678 to 701 bp	CDR3L

(continued)

			from 729 to	o 748 bp	synthetic s	pacer		
5		S: encodes sin ariable domain d	•	ision gene	comprising	monoclonal a	intibody FWP5	1 heavy and kappa
	AAGCT							5
10								
	TCT CAG	GTA CAA	CTG CAG	CAG T	CT GGG	GCT GAG	CTG GTG	44
	Ser Gln	Val Gln	Leu Gln	Gln S	er Gly	Ala Glu	Leu Val	
15	1		5			10		
	AGG CCT	GGG ACT	TCA GTG	AAG C	TG TCC	TGC AAG	GCT TCT	GAT 86
•	Arg Pro	Gly Thr	Ser Val	Lys L	eu Ser	Cys Lys	Ala Ser	Asp
20	15	5		20			25	

 $\varphi_{\hat{G}}^{(k)} \hat{\varphi}_{\hat{A}}$ 

	TAC	ACC	TTC	ACC	AGC	TAC	TGG	ATG	AAC	TGG	GTG	AAG	CAG	AGG	128
_	Tyr	Thr	Phe	Thr	Ser	Tyr	Trp	Met	Asn	Trp	Val	Lys	Gln	Arg	
5			30					35					40		
															150
			CAA												170
10	Pro	Gly	Gln	Gly	Leu	Glu	Trp	Ile		Met	Ile	Asp	Pro		
				45					50					55	
						m».c		C	х <del>пу</del> с	TTTYC	AAG	GAC	AAG	GCC	212
15			GAA												
	Asp	Ser	Glu	Inr	60	Tyr	ASII	GIII	Mec	65	Dy J		2,0		
					60					03					
20	CC A	ጥጥር	ልርጥ	СТА	GAC	AAG	TCC	TCC	AAT	ACA	GCC	TAC	ATG	CAA	254
			Thr												
	70					75					80				
25	. •														
	CTC	AGC	AGC	CTG	ACA	тст	GAG	GAC	TCT	GCG	GTC	TAT	TAC	TGT	296
	Leu	Ser	Ser	Leu	Thr	Ser	Glu	Asp	Ser	Ala	Val	Tyr	Tyr	Cys	
30		85					90					95			
													_		
														TGG	338
35	Ala	Lys	Gly	Gly	Ala	Ser	Gly			Tyr	Phe	Asp		Trp	
			100					105					110		
						0.000		- com	. mcc	· m~		CCT	r ccc	CCT	380
40														GGT	500
	Gly	r Gin	GIY			vai	. 1111	Val	120		. 013	017	011	Gly 125	
				115					120						
45	. ייצרייו	י ככיו	r GGC	GGT	GGC	TCC	GGC	GGT	GGC	GGI	TCT	GAC	ATC	CAG	422
														Gln	
	501		~ <b>_</b> ,		130		-	-	-	135					
50															
	CTO	ACC	CAG	TCI	CCA	TCC	TCA	CTC	TCI	r GC#	TCI	CTC	G GGA	GGC	464
	Let	ı Thr	Gln	Ser	Pro	Sei	. Ser	Lev	seı	. Ala	Ser	Leu	ı Gly	Gly	
<i>55</i>						145	5				150	)			

ann.		GAA	GTC	ACC	ATC	ACT	TGC	AAG	GCA	AGC	CAA	GAC	ATT	AAG	AAG	506
	5	Glu	Val	Thr	Ile	Thr	Cys	Lys	Ala	Ser	Gln	Asp	Ile	Lys	Lys	168
			155					160					165			
																•
		TAT	ATA	GCT	TGG	TAC	CAA	CAC	AAG	CCT	GGA	AAA	AGT	CCT	CGG	548
	10	Tyr	Ile	Ala	Trp	Tyr	Gln	His	Lys	Pro	Gly	Lys	Ser	Pro	Arg	182
				170					175					180		
	15	CTA	CTC	ATA	CAC	TAC	ACA	TCT	GTA	TTA	CAG	CCA	GGC	ATC	CCA	590
		Leu	Leu	Ile	His	Tyr	Thr	Ser	Val	Leu	Gln	Pro	Gly	Ile	Pro	
					185					190					195	
	20															
		TCC	AGG	TTC	AGT	GGA	AGT	GGG	TCT	GGG	AGA	GAT	TAT	TCC	TTC	632
		Ser	Arg	Phe	Ser	Gly	Ser	Gly	Ser	Gly	Arg	Asp	Tyr	Ser	Phe	
	25					200					205					
		AGC	ATC	CAC	AAC	CTG	GAG	CCT	GAA	GAT	ATT	GCA	ACT	TAT	TAT	674
	30	Ser	Ile	His	Asn	Leu	Glu	Pro	Glu	Asp	Ile	Ala	Thr	Tyr	Tyr	
		210					215					220				
;							TAT									716
	35	Cys		His	Tyr	Asp	Tyr		Tyr	Thr	Phe	Gly		Gly	Thr	
			225					230					235			
																- 40
	40					TAG	CTGA'	rca A	AAGC'	I'CTA(	3A					748
		Lys	Leu	Glu	Ile											
				240												
	45															
	S	SEQ ID N	IO:9													
				TYPE:												
	50			LENG1		•										
							: Pseud		s aerug	inosa F	AK					
				ELL CL			URCE:	∟.∞II								
	55 F				·											
	~	FEATUR	RES:	from 1	to 27 b	p	synthet	ic spac	er							

### (continued)

partial exotoxin A sequence corresponding to nucleotide positions 1574 to 1747 bp of the exotoxin A sequence (Gray et al., Proc. Natl. Acad. Sci. USA

5	81: 2645, 1984)	. GCI. GGA
	PROPERTIES: encodes part of the mutated exotoxin A gene from Pseudomonas aeruginosa PAK	
10	AAGCTTAAGG AGATCTGCAT GCTTCTAGAG GGCGGCAGCC TGGCCGCGCT	50
	GACCGCGCAC CAGGCCTGCC ACCTGCCGCT GGAGACTTTC ACCCGTCATC	100
15	GCCAGCCGCG CGGCTGGGAA CAACTGGAGC AGTGCGGCTA TCCGGTGCAG	150
	CGGCTGGTCG CCCTCTACCT GGCGGCGCGA CTGTCATGGA ACCAGGTCGA	200
20	С	201

## SEQ ID NO:10

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SEQUENCE TYPE: nucleotide with corresponding protein

SEQUENCE LENGTH: 2012 bp MOLECULE TYPE: plasmid DNA

ORIGINAL SOURCE ORGANISM: mouse/P.aeruginosa

IMMEDIATE EXPERIMENTAL SOURCE: E.∞Ii

from 29 to 201 bp

NAME OF CELL CLONE: pWW215-5

	FEATURES:	from 1 to 63 bp	ompA signal peptide
35		from 64 to 87 bp	FLAG peptide and enterokinase cleavage site
-		from 97 to 453 bp	FRP5 heavy chain variable domain
		from 454 to 498 bp	15 amino acids linker sequence
		from 499 to 822 bp	FRP5 light chain variable domain
40		from 826 to 1911 bp	exotoxin A gene coding region (coding for amino acids 252 to 613 of the mature exotoxin A)
		from 1912 to 2012 bp	3'non-coding region of the exotoxin A gene

PROPERTIES: Fv heavy chain/light chain variable domain and exotoxin A fusion protein Fv(FRPS)-ETA binding to the c-erbB-2 protein

	ATG	AAA	AAG	ACA	GCT	ATC	GCG	ATT	GCA	GTG	GCA	CTG	GCT	GGT	42
5	Met	Lys	Lys	Thr	Ala	Ile	Ala	Ile	Ala	Val	Ala	Leu	Ala	Gly	
v		-30						-25					-2	20	
	TTC	GCT	ACC	GTT	GCG	CAA	GCT	GAC	TAC	AAG	GAC	GAC	GAT	GAC	84
10	Phe	Ala	Thr	Val	Ala	Gln	Ala	Asp	Tyr	Lys	Asp	Asp	Asp	Asp	
			-15					-10					-5		
15														GAA	126
	Lys	Leu	Ala	Ser	Gln	Val	Gln	Leu	Gln	Gln	Ser	Gly	Pro	Glu	
				1				5					10		
20															

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	CTG	AAG	AAG	ССТ	GGA	GAG	ACA	GTC	AAG	ATC	TCC	TGC	AAG	GCC	168
_	Leu	Lys	Lys	Pro	Gly	Glu	Thr	Val	Lys	Ile	Ser	Cys	Lys	Ala .	
5				15					20					25	
			TAT												210
10	Ser	Gly	Tyr	Pro	Phe	Thr	Asn	Tyr	Gly	Met	Asn	Trp	Val	Lys	
					30					35					
15			CCA												252
	Gln	Ala	Pro	Gly	Gln	Gly	Leu	Lys	Trp	Met	_	Trp	Ile	Asn	
	40					45					50				
20														201	204
														GGA	294
	Thr	Ser	Thr	Gly	Glu	Ser		Phe	Ala	Asp	Asp		ьуs	GIY	
25		55					60					65			
												3.00	000	mam	226
														TAT	336
30	Arg	Phe	Asp	Phe	Ser	Leu	Glu			Ala	ASN	TILL	80	ıyı	
30			70					75					80		
						ama		3.00		CAC	እጥር	CCT	<b>ACA</b>	ጥልጥ	378
														TAT	3.0
35	Leu	Gln	Ile			Leu	rys	Ser	90		Mec	· ALU	****	95	
				85	-				,,						
	mmc	· mca	CCN	እርአ	TICC:	GAG	: ርጥጥ	י ייים	CAC	GGC	TAC	GTT	CCT	TAC	420
40														Tyr	
	Pile	. Cys	AIG	, ALG	100		. ,	-3-		105				_	
					100										
45	ጥረር	: פפר	. СУУ	GGG	: ACC	ACG	GTC	ACC	GTT	TCC	TCT	GGC	GGT	' GGC	462
														Gly	
	110					115					120				
50		•													
	GGT	rTCI	r GGI	GGC	GGI	GGC	TCC	: GGC	GGT	GGC	GGT	TCI	GAC	ATC	504
														Ile	
55	<b>-</b> -	125		_	_		130					135			

,:::...<u>,</u>

	CAG	CTG	ACC	CAG	TCT	CAC	AAA	TTC	CTG	TCC	ACT	TCA	GTA	GGA	546
_	Gln	Leu	Thr	Gln	Ser	His	Lys	Phe	Leu	Ser	Thr	Ser	Val	Gly	
5			140					145					150		
	GAC	AGG	GTC	AGC	ATC	ACC	TGC	AAG	GCC	AGT	CAG	GAT	GTG	TAT	588
10	Asp	Arg	Val	Ser	Ile	Thr	Cys	Lys	Ala	Ser	Gln	Asp	Val	Tyr	
				155					160					165	
15				GCC											630
	Asn	Ala	Val	Ala	Trp	Tyr	Gln	Gln	Lys	Pro	Gly	Gln	Ser	Pro	
					170					175					
20															
														GTC	672
	Lys	Leu	Leu	Ile	Tyr	Ser	Ala	Ser	Ser	Arg	Tyr	Thr	Gly	Val	
25	180					185					190				
2.5															
				TTC											714
	Pro	Ser	Arg	Phe	Thr	Gly	Ser	Gly	Ser	Gly	Pro		Phe	Thr	
30		195					200					205			
														TAT	756
35	Phe	Thr		Ser	Ser	Val	Gln			Asp	Leu	Ala		Tyr	
			210					215					220		
														<b>m</b> cc	700
40														TCG	798
	Phe	Cys	Gln	Gln		Phe	Arg	Thr			Thr	Pne	GIY		
				225					230					235	
45											000	000	100	OMO.	040
														CTG	840
·	Gly	Thr	Lys	Leu			Lys	Ala	Leu			GIY	ser	Leu	
<b>50</b>					240					245					
50				= =				-			~~~	000	cmc	CRO	001
														GAG	882
			Leu	Thr	Ala			Ala	Cys	HIS			ren	GIU	
55	250	)				255					260				

		ACT	TTC	ACC	CGT	CAT	CGC	CAG	CCG	CGC	GGC	TGG	GAA	CAA	CTG	924
et ette		Thr	Phe	Thr	Arg	His	Arg	Gln	Pro	Arg	Gly	Trp	Glu	Gln	Leu	
	5		265					270					275			
		GAG	CAG	TGC	GGC	TAT	CCG	GTG	CAG	CGG	CTG	GTC	GCC	CTC	TAC	966
	10	Glu	Gln	Cys	Gly	Tyr	Pro	Val	Gln	Arg	Leu	Val	Ala	Leu	Tyr	
				280					285					290		
	15	CTG	GCG	GCG	CGA	CTG	TCA	TGG	AAC	CAG	GTC	GAC	CAG	GTG	ATC	1008
		Leu	Ala	Ala	Arg	Leu	Ser	Trp	Asn	Gln	Val	Asp	Gln	Val	Ile	
					295					300					305	
	20,															
	,															1050
		Arg	Asn	Ala	Leu	Ala	Ser	Pro	Gly	Ser	Gly	Gly	Asp	Leu	Gly	
						310					315					
	25															
																1092
		Glu	Ala	Ile	Arg	Glu	Gln	Pro	Glu	Gln	Ala	Arg	Leu	Ala	Leu	
	30	320					325					330				
																1134
i	35	Thr	Leu	Ala	Ala	Ala	Glu		Glu	Arg	Phe	Val		Gln	Gly	
			335					340					345			
																1176
	40															1176
		Thr	Gly		Asp	Glu	Ala	Gly		Ala	Asn	Ala	ASP	Val	vaı	
				350					355					360		
	45						000	000	000	~~m	C 2 2	mcc.	000	ccc	ccc	1210
	40															1218
		Ser	Leu	Thr		Pro	vaı	Ala	Ala		GIU	Cys	ATG	Gly		
					365					370					375	
	50				000	C1C	CCC	CITY C	CEC	CAC	ccc	220	ጥአጥ	ccc	ልርጥ	1260
		-														1260
		Ala	ASP	ser	GIĀ		WIG	neu	red	GIU		ASII	TAT	Pro	1111	
	55					380					385					

	GGC	GCG	GAG	TTC	CTC	GGC	GAC	GGC	GGC	GAC	GTC	AGC	TIC	AGC	1302
	Glv	Ala	Glu	Phe	Leu	Gly	Asp	Gly	Gly	Asp	Val	Ser	Phe	Ser	
5	390					395					400				
	> CC	ccc	ccc	· ACG	CAG	AAC	TGG	ACG	GTG	GAG	CGG	CTG	CTC	CAG	1344
10	ACC	2	03-	Thr	Gln	Asn	Tro	Thr	Val	Glu	Arg	Leu	Leu	Gln	
	Thr			IIIL	GIII	VD	410					415			
		405					140								
					CIII/C	CNG	GAG	CGC	GGC	тат	GTG	TTC	GTC	GGC	1386
15	GCG	CAC	CGG	g Gln		GAG	Clu	λεσ	Glv	Tvr	Val	Phe	Val	Gly	
	Ala	His			Leu	GIU	Giu	425	017	-,-			430		
			42	0				423							
20					_			000	000	CAA	»GC	ልጥሮ	GTC	TTC	1428
	TAC	CAC	GG	C ACC	TTC	CTC	GAA	GCG	33-	CAA	ZO.	Tle	Val	Phe	1428
	Tyr	Hi	s G1	y Thi	: Phe	e Leu	Glu	Ala			per	116	*41	445	
25				435	5				440					1.0	
23												ccc	እጥ <b>Ր</b>	TCC	1470
	GGC	GG	G GT	G CG	C GCC	G CGC	AGC	CAG	GAC	CTC	GAC	310	Tla	ሞፖኮ	1470
	Gly	r G1	y Va	l Ar	g Ala	a Arg	, Ser	Gln	Asp			Ald	116	ııp	
30					450	0				455					
															1512
	CGC	GG	т тт	C TA	T AT	C GCC	GGC	GAT	ccc	GCC	CTG	GCC	TAC	. 666	1512
35	Arg	g Gl	y Ph	ае Ту	r Il	e Ala	a Gly	Asp	Pro	) Ala	1 Leu	Ala	Тух	GIA	•
	46	0				465	5				470	)			
	TA	c GC	C C	AG GA	C CA	G GA	A CCC	GAC	GC	A CG	GGC	CGC	ATC	CGC	1554
40	Τ̈́ν	r Al	a G	ln As	p Gl	n Gl	u Pro	Asp	) Ala	a Ar	g Gly	Arg	g Ile	e Arg	I
		47					480					485	5		
45	λλ	ر در د	T G	CC C1	G CI	G CG	G GT	TA'	r GT	G CC	G CG(	C TC	G AG	CTO	3 1596
	λe	n G	o Iv A	la Le	eu Le	u Ar	g Va	l Ty:	r Va	l Pr	o Arg	g Se	r Se	r Lei	ג
	AS	,, 0.		90			_	49					50	0	
50			-2	. •											
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	PY	O G	TÀ L		71 A	· y · · ·		= •	51					51	5
<i>55</i>				יכ	<i></i>										

	GCG	GCG	GGC	GAG	GTC	GAA	CGG	CTG	ATC	GGC	CAT	CCG	CTG	CCG	1680
	Ala	Ala	Gly	Glu	Val	Glu	Arg	Leu	Ile	Gly	His	Pro	Leu	Pro	
5					520					525					
	CTG	CGC	СПС	GAC	GCC	ATC	ACC	GGC	CCC	GAG	GAG	GAA	GGC	GGG	1722
10		Arg													
		ALG	Dea	ASD	AIG		1111	GIY	110	GIU	540	GIU	OLY	GLY	
	530					535					240				
						0.00	000	maa	000	0.m0	000	010	000	100	1764
15															1764
	Arg	Leu	Glu	Thr	Ile	Leu	_	Trp	Pro	Leu	Ala		Arg	Thr	
		545					550					555			
20															
	GTG	GTG	ATT	CCC	TCG	GCG	ATC	CCC	ACC	GAC	CCG	CGC	AAC	GTC	1806
	Val	Val	Ile	Pro	Ser	Ala	Ile	Pro	Thr	Asp	Pro	Arg	Asn	Val	
25			560					565					570		
25															
	GGC	GGC	GAC	CTC	GAC	CCG	TCC	AGC	ATC	CCC	GAC	AAG	GAA	CAG	1848
	Gly	Gly	Asp	Leu	Asp	Pro	Ser	Ser	Ile	Pro	Asp	Lys	Glu	Gln	
30				575					580				•	585	
	GCG	ATC	AGC	GCC	CTG	CCG	GAC	TAC	GCC	AGC	CAG	CCC	GGC	AAA	1890
35	Ala	Ile	Ser	Ala	Leu	Pro	Asp	Tyr	Ala	Ser	Gln	Pro	Gly	Lys	
					590					595					
	CCG	CCG	CGC	GAG	GAC	CTG	AAG	TAA	CTG	CCGC	GAC (	CGGC	GGC:	ГC	1934
40		Pro													
	600					605	-1 -								
		TCGC	AGG :	AGCC	בפרכי		rcgg	יפררי	r GG(	CAT	ጥልግል	CAGO	ململمك	זייר	1984
45		1000													-,0-
	መሮአ	TGCC	ACC (	ית א חיי	דר כא:	ארדי אר	י ג ג בייו	PTY-							2012
	IGA	1000	noc (	JUMM.	LCGM	al A.	ionn.								2012
50	SEQ ID NO	)· 11													
50	SECTION C	2. 11													
	SEQUI	ENCE T	YPE: n	ucleotic	e with	corresp	onding	protein	1						

SEQUENCE TYPE: nucleotide with corresponding protein SEQUENCE LENGTH: 2012 bp

MOLECULE TYPE: plasmid DNA

ORIGINAL SOURCE ORGANISM: mouse/P.aeruginosa IMMEDIATE EXPERIMENTAL SOURCE: E.∞li

NAME OF CELL CLONE: pWW215-51

5	FEAT	URES:	from from from from from	502 to 826 to	7 bp	F F P P	WP51 I 5 amino WP51 I xotoxin nature e	eptide a heavy o c acids light cha A gene exotoxir	Ind ente Inker s ain vari I coding I A)	erokinas uriable de equence able do g region	domain :e main (codin	g for an		ids 252	to 613 of the
			1 HOIN	1912 (		op 3	non-a	July 16		1110 01	UIUXIII 7	1 yelle			
		OPERT he c-eri			chain/li	ight cha	iin varia	ible dor	nain an	d exoto	xin A fu	sion pr	otein Fv	(FWP5	1)-ETA binding
15	ATG	AAA	AAG	ACA	GCT	ATC	GCG	ATT	GCA	GTG	GCA	CTG	GCT	GGT	42
	Met	Lys	Lys	Thr	Ala	Ile	Ala	Ile	Ala	Val	Ala	Leu	Ala	Gly	
		-30					-25					-20			
20															
	TTC	GCT	ACC	GTT	GCG	CAA	GCT	GAC	TAC	AAG	GAC	GAC	GAT	GAC	84
	Phe	Ala	Thr	Val	Ala	Gln	Ala	Asp	Tyr	Lys	Asp	Asp	Asp	Asp	
25			-15					-10					-5		
	AAG	CTA	GCT	TCT	CAG	GTA	CAA	CTG	CAG	CAG	TCT	GGG	GCT	GAG	126
	Lys	Leu	Ala	Ser	Gln	Val	Gln	Leu	Gln	Gln	Ser	Gly	Ala	Glu	

: *:::.

	CTG	GTG	AGG	CCT	GGG	ACT	TCA	GTG	AAG	CTG	TCC	TGC	AAG	GCT	168
	Leu	Val	Arg	Pro	Gly	Thr	Ser	Val	Lys	Leu	Ser	Cys	Lys	Ala	
5				15					20					25	
	TCT	GAT	TAC	ACC	TTC	ACC	AGC	TAC	TGG	ATG	AAC	TGG	GTG	AAG	210
10			Tyr												
					30					35					
15	CAG	AGG	ССТ	GGA	CAA	GGC	CTT	GAA	TGG	ATT	GGT	ATG	ATT	GAT	252
	Gln	Arg	Pro	Gly	Gln	Gly	Leu	Glu	Trp	Ile	Gly	Met	Ile	Asp	
	40					45					50				
20	ССТ	TCA	GAC	AGT	GAA	ACT	CAA	TAC	AAT	CAA	ATG	TTC	AAG	GAC	294
	Pro	Ser	Asp	Ser	Glu	Thr	Gln	Tyr	Asn	Gln	Met	Phe	Lys	Asp	
		55					60					65			
25															
														TAC	336
	Lys	Ala	Ala	Leu	Thr	Val	Asp	Lys	Ser	Ser	Asn	Thr		Tyr	
30			70					75					80		
															250
														TAT	378
35	Met	Gln	Leu	Ser	Ser	Leu	Thr	Ser			Ser	Ala	Val	Tyr	
				85					90					95	
															420
40														GAT	420
	Tyr	Cys	Ala	Lys			Ala	Ser	Gly			ТУT	Pne	Asp	
					100					105					
4.5			_							· cmm	, mcc	m∕-m		CCTT	462
45														GGT	402
			Gly	Gln	Gly			· vaı	ınr	vaı			GIY	Gly	
	110	)				115					120				
50												CCI	י קער מ	י האר	504
														GAC	304
	Gly			Gly	GIY	GIA			. сту	, сту	GIĀ	135		Asp	
55		125	•				130	,				100	,		

	ATC	CAG	CTG	ACC	CAG	TCT	CCA	TCC	TCA	CTG	TCT	GCA	TCT	CTG	540
	Ile	Gln	Leu	Thr	Gln	Ser	Pro	Ser	Ser	Leu	Ser	Ala	Ser	Leu	
5			140					145					150		
	CCA	GGC	GAA	GTC	ACC	ATC	ACT	TGC	AAG	GCA	AGC	CAA	GAC	ATT	588
10	Gly	Glv	Glu	Val	Thr	Ile	Thr	Cys	Lys	Ala	Ser	Gln	Asp	Ile	
	913	Q ₁	0	155					160					165	
15	AAG	AAG	тат	ATA	GCT	TGG	TAC	CAA	CAC	AAG	CCT	GGA	AAA	AGT	630
	Lvs	Ivs	Tvr	Ile	Ala	Trp	Tyr	Gln	His	Lys	Pro	Gly	Lys	Ser	
	270	_1 -	-3 -		170	_				175					
											-				
20	ССТ	CGG	СТА	CTC	ATA	CAC	TAC	ACA	TCT	GTA	TTA	CAG	CCA	GGC	672
	Pro	Arg	Leu	Leu	Ile	His	Tyr	Thr	Ser	Val	Leu	Gln	Pro	Gly	
	180	3				185					190				
25	100														
	ATC	CCA	TCC	AGG	TTC	AGT	GGA	AGT	GGG	TCT	GGG	AGA	GAT	TAT	714
	Ile	Pro	Ser	Arg	Phe	Ser	Gly	Ser	Gly	Ser	Gly	Arg	Asp	Tyr	
30		195					200					205			
	TCC	TTC	AGC	ATC	CAC	AAC	CTG	GAG	CCT	GAA	GAI	TTA	GCA	ACT	756
35	Ser	Phe	e Ser	Ile	His	Asn	Leu	Glu	Pro	Glu	Asp	Ile	Ala	Thr	
			210					215					220	)	
0.2	TAT	TAT	r TGI	CTA	CAT	'TAT	r GAI	TAT	CTG	TAC	ACG	TTC	GGA	GGG	798
40	Tyr	Туз	c Cys	Lev	n His	Туг	. Asp	Туг	Leu	Тух	Thi	Phe	e Gly	Gly	
	_			225					230					235	
45	GGC	ACC	C AAC	CTC	GAG	TA :	CAAA	GC7	CTA	GAC	GGG	GGG	AGG	CTG	840
	Gly	Th	r Lys	s Lev	ı Glu	ılle	e Lys	ala	Let	ı Glu	ı Gly	, Gl	, Sei	Leu	
					240					245					
50															
	GCC	GC	G CT	G AC	c GCC	CA	C CAC	G GCC	TGO	CAC	CTY	G CC	3 CT	G GAG	882
	Ala	a Al	a Le	u Thi	r Ala	a Hi	s Gl	a Ala	a Cys	, Hi			o Le	ı Glu	
55	250	)				25	5				26	0			

	ACT	TTC	ACC	CGT	CAT	CGC	CAG	CCG	CGC	GGC	TGG	GAA	ÇAA	CTG	924
	Thr	Phe	Thr	Arg	His	Arg	Gln	Pro	Arg	Gly	Trp	Glu	Gln	Leu	
5		265					270					275			
	GAG	CAG	TGC	GGC	TAT	CCG	GTG	CAG	CGG	CTG	GTC	GCC	CTC	TAC	966
10			Cys												
,,			280		•			285					290	-	
	Catc	GCG	GCG	CGA	CTYG	TY	TGG	AAC	CAG	GTYC	GAC	CAG	GTYG	AΤΥ	1008
15			Ala												
	Deu	AIG	AIG	_	beu	361	пр	ASII		vai	rop	Gin	Val	305	
				295					300					303	
20	666		000	~m^	000	200	~~~	<b>~</b>	100	666	000	C) C	CmC	ccc	1050
															1050
	Arg	ASII	Ala	Leu		ser	PIO	GIÀ	ser		GTĀ	ASD	Leu	GIY	
					310					315					
25															
															1092
		Ala	Ile	Arg	Glu	Gln	Pro	Glu	Gln	Ala	_	Leu	Ala	Leu	
30	320					325					330				
	ACC	CTG	GCC	GCC	GCC	GAG	AGC	GAG	CGC	TTC	GTC	CGG	CAG	GGC	1134
35	Thr	Leu	Ala	Ala	Ala	Glu	Ser	Glu	Arg	Phe	Val	Arg	Gln	Gly	
		335					340					345			
	ACC	GGC	AAC	GAC	GAG	GCC	GGC	GCG	GCC	AAC	GCC	GAC	GTG	GTG	1176
40	Thr	Gly	Asn	Asp	Glu	Ala	Gly	Ala	Ala	Asn	Ala	Asp	Val	Val	
			350					355					360		
45	AGC	CTG	ACC	TGC	CCG	GTC	GCC	GCC	GGT	GAA	TGC	GCG	GGC	CCG	1218
	Ser	Leu	Thr	Cys	Pro	Val	Ala	Ala	Gly	Glu	Cys	Ala	Gly	Pro	
				365					370					375	
50															
50	GCG	GAC	AGC	GGC	GAC	GCC	CTG	CTG	GAG	CGC	AAC	TAT	ccc	ACT	1260
			Ser												
			<b></b>		380					385					
55					555					303					

	GGC	GCG	GAG	TTC	CTC	GGC	GAC	GGC	GGC	GAC	GTC	AGC	TTC	AGC	1302
	Gly	Ala	Glu	Phe	Leu	Gly	Asp	Gly	Gly	Asp	Val	Ser	Phe	Ser	
5	390					395					400				
	ACC	CGC	GGC	ACG	CAG	AAC	TGG	ACG	GTG	GAG	CGG	CTG	CTC	CAG	1344
10	Thr	Arg	Gly	Thr	Gln	Asn	Trp	Thr	Val	Glu	Arg	Leu	Leu	Gln	448
		405					410				•	415			
15	GCG	CAC	CGC	CAA	CTG	GAG	GAG	CGC	GGC	TAT	GTG	TTC	GTC	GGC	1386
	Ala	His	Arg	Gln	Leu	Glu	Glu	Arg	Gly	Tyr	Val	Phe	Val	Gly	
			420					425					430		
20															
20	TAC	CAC	GGC	ACC	TTC	CTC	GAA	GCG	GCG	CAA	AGC	ATC	GTC	TTC	1428
	Tyr	His	Gly	Thr	Phe	Leu	Glu	Ala	Ala	Gln	Ser	Ile	Val	Phe	
				435					440					445	
25															•
	GGC	GGG	GTG	CGC	GCG	CGC	AGC	CAG	GAC	CTC	GAC	GCG	ATC	TGG	1470
	Gly	Gly	val	Arg	Ala	Arg	Ser	Gln	Asp	Leu	Asp	Ala	Ile	Trp	
30					450					455					
													_		
	CGC	GG1	TTC	TAT	ATC	GCC	GGC	GAT	, CCC	GCG	CTG	GCC	TAC	GGC	1512
35	Arg	Gly	, Phe	Tyr	Ile	Ala	Gly	Asp	Pro	Ala	Leu	Ala	Tyr	Gly	
	460	)				465	,				470				
															3554
40	TAC	GC	CAG	GAC	CAG	GAA	CCC	GAC	GCA	A CGC	GGC	CGG	ATC	CGC	1554
40	Ту	. Ala	a Glr	n Asp	Gln	Glu	Pro	) Asr	) Alā	a Arg	g Gly			Arg	
		47	5				480	)				485	•		
															1500
45	AAG	G GG	r GCC	CTC	CTC	CGG	GTC	TAT	r GTC	G CCC	G CGC	TCG	AGC	CIG	1596
	Ası	n Gl	y Ala	a Lei	ı Lev	Arg	y Val	LTy	r Val	l Pro	o Arg	, Sei		Leu	•
			490	0				49	5				500	,	
50															1 626
	CC	G GG	C TI	C TAC	CGC	: ACC	C AGO	CTY	G AC	CT	G GCC	; GC(	5 CC(	GAG	1638
	Pr	o Gl	y Ph	е Ту	r Arg	Th	r Se	r Le			u Ala	A AL	a PTC	Glu 515	
55				50	5				51	0				515	,

5	GCG GCG GGC GAG GTC GAA CGG CTG ATC GGC CAT CCG CTG CCG 1680 Ala Ala Gly Glu Val Glu Arg Leu Ile Gly His Pro Leu Pro 520 525
10	CTG CGC CTG GAC GCC ATC ACC GGC CCC GAG GAG GAA GGC GGG 1722  Leu Arg Leu Asp Ala Ile Thr Gly Pro Glu Glu Glu Gly Gly  530 540
15	CGC CTG GAG ACC ATT CTC GGC TGG CCG CTG GCC GAG CGC ACC 1764  Arg Leu Glu Thr Ile Leu Gly Trp Pro Leu Ala Glu Arg Thr  545 550 555
20	GTG GTG ATT CCC TCG GCG ATC CCC ACC GAC CCG CGC AAC GTC 1806 Val Val Ile Pro Ser Ala Ile Pro Thr Asp Pro Arg Asn Val 560 565 570
25 30	GGC GGC GAC CTC GAC CCG TCC AGC ATC CCC GAC AAG GAA CAG 1848 Gly Gly Asp Leu Asp Pro Ser Ser Ile Pro Asp Lys Glu Gln 575 580 585
35	GCG ATC AGC GCC CTG CCG GAC TAC GCC AGC CAG CCC GGC AAA 1890 Ala Ile Ser Ala Leu Pro Asp Tyr Ala Ser Gln Pro Gly Lys 590 595
40	CCG CCG CGC GAG GAC CTG AAG TAA CTGCCGCGAC CGGCCGGCTC 1934 Pro Pro Arg Glu Asp Leu Lys 600 605
45	CCTTCGCAGG AGCCGGCCTT CTCGGGGCCT GGCCATACAT CAGGTTTTCC 1984
50	TGATGCCAGC CCAATCGAAT ATGAATTC 2012 Claims

- Claims for the following contracting States : AT, BE, CH, LI, DE, DK, FR, GB, GR, IT, LU, NL, PT, SE
  - A fusion protein comprising a single chain recombinant antibody directed to the extracellular domain of the growth factor receptor c-erbB-2 comprising a heavy chain variable domain and a light chain variable domain of a mono-

clonal antibody which domains are linked by a polypeptide spacer group and an effector molecule, and optionally comprising a peptide facilitating purification, a cleavage site and a peptide spacer.

2. A fusion protein according to claim 1 wherein the heavy chain variable domain comprises a polypeptide of the amino acid sequence 2 to 120 of SEQ ID NO:4, wherein optionally 1, 2, 3 or 4 single amino acids within the amino acid sequences 2 to 31 (FR₁), 37 to 50 (FR₂), 68 to 99 (FR₃), and/or 110 to 120 (FR₄) are replaced by other amino acids or deleted, and wherein the amino acid Cys may be in the oxidized state forming S-S-bridges.

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- A fusion protein according to claim 1 wherein the heavy chain variable domain comprises a polypeptide of the amino acid sequence 2 to 120 of SEQ ID NO:4, wherein the amino acid Cys may be in the oxidized state forming S-S-bridges.
  - 4. A fusion protein according to claim 1 wherein the light chain variable domain comprises a polypeptide of the amino acid sequence 136 to 241 of SEQ ID NO:4, wherein optionally 1, 2, 3 or 4 single amino acids within the amino acid sequences 136 to 158 (FR₆), 170 to 184 (FR₇), 192 to 223 (FR₈), and/or 233 to 241 (FR₉) are replaced by other amino acids or deleted, and wherein the amino acid Cys may be in the oxidized state forming S-S-bridges.
  - A fusion protein according to claim 4 wherein the light chain variable domain comprises a polypeptide of the amino acid sequence 136 to 241 of SEQ ID NO:4, wherein the amino acid Cys may be in the oxidized state forming S-S-bridges.
  - 6. A fusion protein according to claim 1 wherein the heavy chain variable domain comprises a polypeptide of the amino acid sequence 2 to 121 of SEQ ID NO:8, wherein optionally 1, 2, 3 or 4 single amino acids within the amino acid sequences 2 to 31 (FR₁), 37 to 50 (FR₂), 68 to 99 (FR₃), and/or 111 to 121 (FR₄) are replaced by other amino acids or deleted, and wherein the amino acid Cys may be in the oxidized state forming S-S-bridges.
  - 7. A fusion protein according to claim 6 wherein the heavy chain variable domain comprises a polypeptide of the amino acid sequence 2 to 121 of SEQ ID NO:8, wherein the amino acid Cys may be in the oxidized state forming S-S-bridges.
  - 8. A fusion protein according to claim 6 wherein the light chain variable domain comprises a polypeptide of the amino acid sequence 137 to 241 of SEQ ID NO:8, wherein optionally one or more single amino acids within the amino acid sequences 137 to 159 (FR₆), 171 to 185 (FR₇), 193 to 224 (FR₈), and/or 233 to 241 (FR₈) are replaced by other amino acids or deleted, and wherein the amino acid Cys may be in the oxidized state forming S-S-bridges.
  - A fusion protein according to claim 8 wherein the light chain variable domain comprises a polypeptide of the amino acid sequence 137 to 241 of SEQ ID NO:8, wherein the amino acid Cys may be in the oxidized state forming S-S-bridges.
- 40 10. A fusion protein according to any of claims 1 to 9 wherein the effector molecule is an enzyme or a biologically active variant thereof.
  - A fusion protein according to claim 10 wherein the enzyme is alkaline phosphatase or a biologically active variant thereof.
  - 12. A fusion protein according to any of claims 1 to 9 wherein the effector molecule is a toxin or a biologically active variant thereof.
  - 13. A fusion protein according to claim 12 wherein the effector molecule is Pseudomonas exotoxin or a biologically active variant therof.
  - 14. A fusion protein according to claim 1 wherein the heavy chain variable domain and the light chain variable domain are derived from a mouse monoclonal antibody selected from the group consisting of FRPS, FSP16, FWP51 and FSP77 deposited under the Budapest Treaty on November 21, 1990 at the European Collection of Animal Cell Cultures (ECACC) in Porton Down, Salisbury, UK, under accession numbers 90112115, 90112116, 90112117, and 90112118, respectively.
  - 15. A fusion protein according to claim 1 wherein the heavy chain variable domain and the light chain variable domain

are derived from the mouse monoclonal antibody FRP5.

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- 16. A fusion protein according to claim 1 wherein the heavy chain variable domain and the light chain variable domain are derived from the mouse monoclonal antibody SWP51.
- A fusion protein designated Fv(FRP5)-phoA according to claim 1 comprising a polypeptide of the amino acid sequence 2 to 690 of SEQ ID NO:5.
- A fusion protein designated Fv(FRP5)-ETA according to claim 1 comprising a polypeptide of the amino acid sequence 2 to 606 of SEQ ID NO:10.
  - A fusion protein designated Fv(FWP51)-ETA according to claim 1 comprising a polypeptide of the amino acid sequence 2 to 606 of SEQ ID NO: 11.
- 15 20. A recombinant DNA comprising an insert coding for a fusion protein according to any of claims 1 to 19.
  - 21. A recombinant DNA according to claim 20 comprising an insert coding for a heavy chain murine variable domain of a monoclonal antibody selected from the group consisting of antibodies FRP5, FSP16, FSP77 and FWP51 deposited under the Budapest Treaty on November 21, 1990 at the European Collection of Animal Cell Cultures (ECACC) in Porton Down, Salisbury, UK, under accession numbers 90112115, 90112116, 90112117, and 90112118, respectively, or coding for an amino acid sequence homologous to said heavy chain variable domain.
  - 22. A recombinant DNA according to claim 21 comprising an insert coding for a light chain murine variable domain of a monoclonal antibody selected from the group consisting of antibodies FRP5, FSP16, FSP77 and FWP51 deposited under the Budapest Treaty on November 21, 1990 at the European Collection of Animal Cell Cultures (ECACC) in Porton Down, Salisbury, UK, under accession numbers 90112115, 90112116, 90112117, and 90112118, respectively, or coding for an amino acid sequence homologous to said light chain variable domain.
- 23. A recombinant DNA according to claim 20 which is a hybrid vector further comprising an origin of replication or an autonomously replicating sequence, one or more dominant marker sequences and, optionally, expression control sequences, signal sequences and additional restriction sites.
  - 24. A hybrid vector according to claim 23 comprising a Simian virus promoter and the mouse tg H or L chain enhancer.
- 25. A process for the preparation of a DNA according to claim 20 comprising the steps of
  - a) preparing murine DNA from the genome of a suitable hybridoma cell line and selecting the desired DNA coding for the variable heavy and/or light chain domains of the antibody with the desired specificity,
  - b) preparing DNA coding for the desired signal sequence and preparing DNA coding for an effector molecule,
  - c) synthesizing DNA coding for the desired spacer group by chemical methods,
  - d) constructing recombinant genes encoding the fusion proteins by incorporating the DNA of step a) and c), and optionally b), into appropriate hybrid vectors,
  - e) transferring the obtained hybrid vectors into a recipient host cell or retrieving the DNA coding for the recombinant genes and transferring the unlinked DNA into a recipient host cell,
  - f) selecting and culturing the transformed host cell, and
  - g) optionally isolating the desired DNA.
  - 26. A host cell transformed with a recombinant DNA according to claim 25.
- 50 27. A host cell according to claim 26 which is a cell of a strain of E. coli.
  - 28. A process for the preparation of a transformed host cell according to claim 26 wherein suitable recipient cells are transformed with a hybrid vector comprising a DNA insert according to claim 20, an origin of replication or an autonomously replicating sequence, one or more dominant marker sequences and, optionally, expression control sequences, signal sequences and additional restriction sites, and the transformed cells are selected.
  - Use of a fusion protein according to claim 1 for the qualitative and quantitative determination of the growth factor receptor c-erbB-2.

- 30. Use according to claim 29 comprising immunostaining of tissue sections with a solution containing the fusion protein comprising a detectable enzyme.
- A test kit for the qualitative and quantitative determination of c-erbB-2 protein comprising a fusion protein according to claim 1.
- 32. A fusion protein according to claim 1 for use in the treatment of the human or animal body.
- 33. A pharmaceutical composition for treating tumors over-expressing the growth factor receptor c-erbB-2 comprising a therapeutically effective amount of a fusion protein according to claim 1 and a pharmaceutically acceptable carrier.
- 34. The use of a fusion protein according to claim 1 for the manufacture of a pharmaceutical preparation.

### 5 Claims for the following contracting State: ES

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- Process for the preparation of a fusion protein comprising a single chain recombinant antibody directed to the
  extracellular domain of the growth factor receptor c-erbB-2 comprising a heavy chain variable domain and a light
  chain variable domain of a monoclonal antibody which are linked by a polypeptide spacer, and an effector group,
  characterised in that cells producing such an antibody are multiplied in vitro or in vivo and, when required, the
  obtained antibody is isolated.
- 2. Process according to claim 1 for the preparation of a fusion protein wherein the heavy chain variable domain comprises a polypeptide of the amino acid sequence 2 to 120 of SEQ ID NO:4, wherein optionally 1, 2, 3 or 4 single amino acids within the amino acid sequences 2 to 31 (FR₁), 37 to 50 (FR₂), 68 to 99 (FR₃), and/or 110 to 120 (FR₄) are replaced by other amino acids or deleted, and wherein the amino acid Cys may be in the oxidised state forming S-S-bridges.
- 3. Process according to claim 1 for the preparation of a fusion protein wherein the heavy chain variable domain comprises a polypeptide of the amino acid sequence 2 to 120 of SEQ ID NO:4, wherein the amino acid Cys may be in the oxidised state forming S-S-bridges.
  - 4. Process according to claim 1 for the preparation of a fusion protein wherein the light chain variable domain comprises a polypeptide of the amino acid sequence 136 to 241 of SEQ ID NO:4, wherein optionally 1, 2, 3 or 4 single amino acids within the amino acid sequences 136 to 158 (FR₆), 170 to 184 (FR₇), 192 to 223 (FR₈), and/or 233 to 241 (FR₉) are replaced by other amino acids or deleted, and wherein the amino acid Cys may be in the oxidised state forming S-S-bridges.
- 5. A process according to claim 4 for the manufacture of a fusion protein wherein the light chain variable domain comprises a polypeptide of the amino acid sequence 136 to 241 of SEQ ID NO:4, wherein the amino acid Cys may be in the oxidised state forming S-S-bridges.
  - 6. A process according to claim 1 for the manufacture of a fusion protein wherein the heavy chain variable domain comprises a polypeptide of the amino acid sequence 2 to 121 of SEQ ID NO:8, wherein optionally 1, 2, 3 or 4 single amino acids within the amino acid sequences 2 to 31 (FR₁), 37 to 50 (FR₂), 68 to 99 (FR₃), and/or 111 to 121 (FR₄) are replaced by other amino acids or deleted, and wherein the amino acid Cys may be in the oxidised state forming S-S-bridges.
- 7. A process according to claim 6 for the manufacture of a fusion protein wherein the heavy chain variable domain comprises a polypeptide of the amino acid sequence 2 to 121 of SEQ ID NO:8, wherein the amino acid Cys may be in the oxidised state forming S-S-bridges.
  - 8. A process according to claim 6 for the manufacture of a fusion protein wherein the light chain variable domain comprises a polypeptide of the amino acid sequence 137 to 241 of SEQ ID NO:8, wherein optionally one or more single amino acids within the amino acid sequences 137 to 159 (FR₆), 171 to 185 (FR₇), 193 to 224 (FR₈), and/or 233 to 241 (FR₈) are replaced by other amino acids or deleted, and wherein the amino acid Cys may be in the oxidised state forming S-S-bridges.

- A process according to claim 8 for the manufacture of a fusion protein wherein the light chain variable domain comprises a polypeptide of the amino acid sequence 137 to 241 of SEQ ID NO:8, wherein the amino acid Cys may be in the oxidised state forming S-S-bridges.
- 5 10. A process according to claim 1 for the manufacture of a fusion protein wherein the effector molecule is an enzyme or a biologically active variant thereof.
  - 11. A process according to claim 10 wherein the enzyme is alkaline phosphatase or a biologically active variant thereof.
- 9 12. A process according to claim 10 wherein the effector molecule is a toxin or a biologically active variant thereof.
  - A process according to claim 12 wherein the effector molecule is Pseudomonas exotoxin or a biologically active variant thereof.
- 14. A process according to claim 1 for the manufacture of a fusion protein wherein the heavy chain variable domain and the light chain variable domain are derived from a mouse monoclonal antibody selected from the group consisting of FRP5, FSP16, FWP51 and FSP77 deposited under the Budapest Treaty on November 21, 1990 at the European Collection of Animal Cell Cultures (ECACC) in Porton Down, Salisbury, UK, under accession numbers 90112115, 90112116, 90112117, and 90112118, respectively.

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- 15. A process according to claim 1 wherein the heavy chain variable domain and the light chain variable domain are derived from the mouse monoclonal antibody FRPS.
- 16. A process according to claim 1 wherein the heavy chain variable domain and the light chain variable domain are derived from the mouse monoclonal antibody FWP51.
- 17. A process according to claim 1 for the manufacture of a fusion protein designated Fv(FRP5)-phoA comprising a polypeptide of the amino acid sequence 2 to 690 of SEQ ID NO:5.
- 30 18. A process according to claim 1 for the manufacture of a fusion protein designated Fv(FRP5)-ETA 1 comprising a polypeptide of the amino acid sequence 2 to 606 of SEQ ID NO:10.
  - A process according to claim 1 for the manufacture of a fusion protein designated Fv(FWP51)-ETA comprising a
    polypeptide of the amino acid sequence 2 to 606 of SEQ ID NO:11.
  - 20. A process according to claim 1 wherein the cells expressing the fusion protein are transformed with a recombinant DNA sequence encoding said fusion protein.
  - 21. A process according to claim 20 wherein the recombinant DNA sequence comprises an insert coding for a heavy chain murine variable domain of a monoclonal antibody selected from the group consisting of antibodies FRP5, FSP16, FSP77 and FWP51 deposited under the Budapest Treaty on November 21, 1990 at the European Collection of Animal Cell Cultures (ECACC) in Porton Down, Salisbury, UK, under accession numbers 90112115, 90112116, 90112117, and 90112118, respectively, or coding for an amino acid sequence homologous to said heavy chain variable domain.
    - 22. A process according to claim 20 wherein the recombinant DNA sequence comprises an insert coding for a light chain murine variable domain of a monoclonal antibody selected from the group consisting of antibodies FRP5, FSP16, FSP77 and FWP51 deposited under the Budapest Treaty on November 21, 1990 at the European Collection of Animal Cell Cultures (ECACC) in Porton Down, Salisbury, UK, under accession numbers 90112115, 90112116, 90112117, and 90112118, respectively, or coding for an amino acid sequence homologous to said light chain variable domain.
    - 23. A process according to claim 20 wherein the recombinant DNA sequence is comprised in a hybrid vector further comprising an origin of replication or an autonomously replicating sequence, one or more dominant marker sequences and, optionally, expression control sequences, signal sequences and additional restriction sites.
    - 24. A process according to claim 23 wherein the hybrid vector further comprises a Simian virus promoter and the mouse Ig H or L chain enhancer.

- 25. A process for the preparation of a recombinant DNA sequence according to claim 20 comprising the steps of
  - a) preparing murine DNA from the genome of a suitable hybridoma cell line and selecting the desired DNA coding for the variable heavy and/or light chain domains of the antibody with the desired specificity,
  - b) preparing DNA coding for the desired signal sequence and/or preparing DNA coding for an effector molecule,
  - c) synthesising DNA coding for the desired spacer group by chemical methods,
  - d) constructing recombinant genes encoding the recombinant antibodies by incorporating the DNA of step a) and, optionally, b) and/or c) into appropriate hybrid vectors,
  - e) transferring the obtained hybrid vectors into a recipient host cell or retrieving the DNA coding for the recombinant genes and transferring the unlinked DNA into a recipient host cell,
  - f) selecting and culturing the transformed host cell, and
  - g) optionally isolating the desired DNA.
- 26. A process for the preparation of a host cell transformed with a recombinant DNA sequence according to claim 20 wherein suitable recipient cells are transformed with a hybrid vector according to claim 23, an origin of replication or an autonomously replicating sequence, one or more dominant marker sequences and, optionally, expression control sequences, signal sequences and additional restriction sites, and the transformed cells are selected.

### 20 Patentansprüche

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### Patentansprüche für folgende Vertragsstaaten : AT, BE, CH, LI, DE, DK, FR, GB, GR, IT, LU, NL, PT, SE

- 25 1. Fusionsprotein, das einen rekombinanten Einketten-Antik\u00f6rper umfa\u00e4t, der gegen die extrazellul\u00e4re Dom\u00e4ne des Wachstumsfaktor-Rezeptors c-erbB-2 gerichtet ist, welcher eine variable Dom\u00e4ne einer schweren Kette und eine variable Dom\u00e4ne einer leichten Kette eines monoklonalen Antik\u00f6rpers, wobei die Dom\u00e4nen durch eine Polypeptid-Spacergruppe verbunden sind, und ein Effektormolek\u00fcl umfa\u00e4t, und gegebenenfalls ein Peptid, das die Reinigung erleichtert, eine Spaltstelle und einen Peptidspacer umfa\u00e4t.
  - 2. Fusionsprotein nach Anspruch 1, worin die variable Dornäne der schweren Kette ein Polypeptid der Aminosäure-sequenz 2-120 der SEQ ID Nr.4 umfaßt, worin gegebenenfalls 1, 2, 3 oder 4 einzelne Aminosäuren in den Aminosäuresequenzen 2-31 (FR₁), 37-50 (FR₂), 68-99 (FR₃) und/oder 110-120 (FR₄), durch andere Aminosäuren ersetzt sind oder entfernt wurden, und worin die Aminosäure Cys in dem oxidierten Zustand unter Bildung von S-Brücken vorliegen kann.
  - Fusionsprotein nach Anspruch 1, worin die variable Dom\u00e4ne der schweren Kette ein Polypeptid der Aminos\u00e4uresequenz 2-120 der SEQ ID Nr.4 umfa\u00dft, worin die Aminos\u00e4ure Cys in dem oxidierten Zustand unter Bildung von S-S-Br\u00fccken vorliegen kann.
  - 4. Fusionsprotein nach Anspruch 1, worin die variable Domâne der leichten Kette ein Polypeptid der Aminosäuresequenz 136-241 der SEQ ID Nr.4 umfaßt, worin gegebenenfalls 1, 2, 3 oder 4 einzelne Aminosäuren in den Aminosäuresequenzen 136-158 (FR₆), 170-184 (FR₇), 192-223 (FR₈), und/oder 233-241 (FR₉) durch andere Aminosäuren ersetzt sind, oder entfernt wurden, und worin die Aminosäure Cys in dem oxidierten Zustand unter Bildung von S-S-Brücken vorliegen kann.
  - Fusionsprotein nach Anspruch 4, worin die variable Domäne der leichten Kette ein Polypeptid der Aminosäuresequenz 136-241 der SEQ ID Nr.4 umfaßt, worin die Aminosäure Cys in dem oxidierten Zustand unter Bildung von S-S-Brücken vorliegen kann.
  - 6. Fusionsprotein nach Anspruch 1, worin die variable Domäne der schweren Kette ein Polypeptid der Aminosäure-sequenz 2-121 der SEQ ID Nr.8 umfaßt, worin gegebenenfalls 1, 2, 3 oder 4 einzelne Aminosäuren in den Aminosäuresequenzen 2-31 (FR₁), 37-50 (FR₂), 68-99 (FR₃) und/oder 111-121 (FR₄) durch andere Aminosäuren ersetzt sind oder entfernt wurden, und worin die Aminosäure Cys in dem oxidierten Zustand unter Bildung von S-S-Brücken vorliegen kann.
  - Fusionsprotein nach Anspruch 6, worin die variable Dom\u00e4ne der schweren Kette ein Polypeptid der Aminos\u00e4uresequenz 2-121 der SEQ ID Nr.8 umla\u00dft, worin die Aminos\u00e4ure Cys in dem oxidierten Zustand unter Bildung von

### S-S- Brücken vorliegen kann

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- 8. Fusionsprotein nach Anspruch 6, worin die variable Domäne der leichten Kette ein Polypeptid der Aminosäure-sequenz 137-241 der SEQ ID Nr.8 umfaßt, worin gegebenenfalls eine oder mehrere einzelne Aminosäuren in den Aminosäuresequenzen 137-159 (FR₆); 171-185 (FR₇), 193-224 (FR₈) und/oder 233-241 (FR₉) durch andere Aminosäure ersetzt sind oder entfernt wurden, und worin die Aminosäure Cys in dem oxidierten Zustand unter Bildung von S-S-Brücken vorliegen kann.
- Fusionsprotein nach Anspruch 8, worin die variable Dom\u00e4ne der leichten Kette ein Polypeptid der Aminos\u00e4urere sequenz 137-241 der SEQ ID Nr.8 umfa\u00d8t, worin die Aminos\u00e4ure Cys in dem oxidierten Zustand unter Bildung
  von S-S-Br\u00fccken vorliegen kann
  - 10. Fusionsprotein nach einem der Ansprüche 1 bis 9, worin das Effektormolekül ein Enzym oder eine biologisch aktive Variante davon ist.
  - Fusionsprotein nach Anspruch 10, worin das Enzym alkalische Phosphatase oder eine biologisch aktive Variante davon ist.
  - Fusionsprotein nach einem der Ansprüche 1 bis 9, worin das Effektormolekül ein Toxin oder eine biologisch aktive Variante davon ist.
    - Fusionsprotein nach Anspruch 12, worin das Effektormolekül Pseudomonas Exotoxin oder eine biologisch aktive Variante davon ist.
- 25 14. Fusionsprotein nach Anspruch 1, worin die variable Dom\u00e4ne der schweren Kette und die variable Dom\u00e4ne der leichten Kette von einem monoklonalen Maus-Antik\u00f6rper abgeleitet ist, der ausgew\u00e4hlt ist aus der Gruppe bestehend aus FRP5, FSP16, FWP51 und FSP77, die die gem\u00e4\u00e4 dem Budapester Vertrag am 21. November 1990 bei der European Collection of Animal Cell Cultures (ECACC) in Porton Down, Salisbury, GB, unter den Hinterlegungsnummern 90112115, 90112116, 90112117 bzw. 90112118 hinterlegt wurden.
  - 15. Fusionsprotein nach Anspruch 1, worin die variable Dom\u00e4ne der schweren Kette und die variable Dom\u00e4ne der leichten Kette von dem monoklonalen Maus-Antik\u00f6rper FRP5 abgeleitet sind.
  - 16. Fusionsprotein nach Anspruch 1, worin die variable Dom\u00e4ne der schweren Kette und die variable Dom\u00e4ne der leichten Kette von dem monoklonalen Maus-Antik\u00f6rper FWP51 abgeleitet sind.
  - Fusionsprotein, das als Fv(FRP5)-phoA bezeichnet wird, nach Anspruch 1, welches ein Polypeptid der Aminosäuresequenz 2-690 der SEQ ID Nr.5 umfaßt.
- 40 18. Fusionsprotein, das als Fv(FRP5)-ETA bezeichnet wird, nach Anspruch 1, welches ein Polypeptid der Aminosäuresequenz 2-606 der SEQ ID Nr.10 umfaßt.
  - Fusionsprotein, das als Fv(FWP51)-ETA bezeichnet wird, nach Anspruch 1, welches ein Polypeptid der Aminosäuresequenz 2-606 der SEQ ID Nr.II umfaßt.
  - 20. Rekombinante DNA, welche ein Insert umfaßt, das ein Fusionsprotein nach einem der Ansprüche 1 bis 19 codiert.
  - 21. Rekombinante DNA nach Anspruch 20, die ein Insert umfaßt, das eine variable Domäne einer schweren Kette der Maus eines monoklonalen Antikörpers codiert, der ausgewählt ist aus der Gruppe bestehend aus den Antikörpern FRP5, FSP16, FSP77 und FWP51, die gemäß dem Budapester Vertrag am 21. November 1990 bei der European Collection of Animal Cell Cultures (ECACC) in Porton Down, Salisbury, GB, unter den Hinterlegungsnummem 90112115, 90112116, 90112117 bzw. 90112118 hinterlegt wurden, oder eine Aminosäuresequenz codiert, die zu der variablen Domäne der schweren Kette homolog ist.
- 22. Rekombinante DNA nach Anspruch 21, die ein Insert umfaßt, das eine variable Domäne einer leichten Kette der Maus eines monoklonalen Antikörpers codiert, der ausgewählt ist aus der Gruppe bestehend aus den Antikörpern FRP5, FSP16, FSP77 und FWP51, die gemäß dem Budapester Vertrag am 21. November 1990 bei der European Collection of Animal Cell Cultures (ECACC) in Porton Down, Salisbury, GB, unter den Hinterlegungsnummern

90112115, 90112116, 90112117 bzw. 90112118 hinterlegt wurden, oder eine Aminosäuresequenz codiert, die zu der variablen Domäne der leichten Kette homolog ist.

- 23. Rekombinante DNA nach Anspruch 20, die ein Hybridvektor ist, der weiter einen Ursprung der Replikation oder eine autonom replizierende Sequenz, einen oder mehrere dominante Markersequenzen, und gegebenenfalls Expressionskontrollsequenzen, Signalsequenzen und zusätzliche Restriktionschnittstellen enthält.
- Hybridvektor nach Anspruch 23, der einen Aflen-Virus-Promotor und den Enhancer der Maus Ig H- oder L-Kette umfaßt
- 25. Verfahren zur Herstellung einer DNA nach Anspruch 20, welches die folgenden Schritte umfaßt:
  - a) Herstellen von Maus-DNA aus dem Genom einer geeigneten Hybridomzellinie und Auswählen der gewünschten DNA, die die variablen Domänen der schweren und/oder leichten Kette des Antikörpers mit der gewünschten Spezifität codiert,
  - b) Herstellen von DNA, die die gewünschte Signalsequenz codiert und Herstellen von DNA, die ein Effektormolekül codiert,
  - c) Synthetisieren von DNA mittels chemischer Methoden, die die gewünschte Spacergruppe codiert,
  - d) Bilden rekombinanter Gene, die die Fusionsproteine codieren, indem die DNA aus Schritt a) und c) und gegebenenfalls b) in geeignete Hybridvektoren eingebaut werden,
  - e) Transferieren der erhaltenen Hybridvektoren in eine aufnehmende Wirtszelle oder Wiedergewinnen der DNA, die die rekombinanten Gene codiert und Transferieren der nicht-verknüpften DNA in eine aufnehmende Wirtszelle.
  - f) Auswählen und Züchten der transformierten Wirtszelle, und
  - g) gegebenenfalls Isolieren der gewünschten DNA.

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- 26. Wirtszelle, die mit einer rekombinanten DNA nach Anspruch 25 transformiert ist.
- 27. Wirtszelle nach Anspruch 26, die eine Zelle eines E.coli-Stamms ist.
- 28. Verfahren zur Herstellung einer transformierten Wirtszelle nach Anspruch 26, wobei geeignete aufnehmende Zellen mit einem Hybridvektor transformiert werden, der ein DNA-Insert nach Anspruch 20, einen Ursprung der Replikation oder eine autonom replizierende Sequenz, einen oder mehrere dominante Markersequenzen und gegebenenfalls Expressionskontrollsequenzen, Signalsequenzen und zusätzlich Restriktionsschnittstellen umfaßt, und die transformierten Zellen ausgewählt werden.
- Verwendung eines Fusionsproteins nach Anspruch 1 zur qualitativen und quantitativen Bestimmung des Wachstumsfaktor-Rezeptors c-erbB-2.
- 40 30. Verwendung nach Anspruch 29, welches Immunfärben von Gewebeschnitten mit einer Lösung umfaßt, die das Fusionsprotein, welches ein nachweisbares Enzym umfaßt, enthält.
  - Test-Kit zur qualitativen und quantitativen Bestimmung des c-erbB-2-Proteins, welcher ein Fusionsprotein nach Anspruch 1 umfaßt.
  - 32. Fusionsprotein nach Anspruch 1 zur Verwendung bei der Behandlung des menschlichen oder tierischen Körpers.
  - 33. Pharmazeutische Zusammensetzung zur Behandlung von Tumoren, die den Wachstumsfaktor-Rezeptor c-erbB-2 überexprimieren, welche eine therapeutisch wirksame Menge eines Fusionsproteins nach Anspruch 1 und einen pharmazeutisch verträglichen Träger umfaßt.
  - 34. Verwendung eines Fusionsproteins nach Anspruch 1 zur Herstellung einer pharmazeutischen Zubereitung.

## 55 Patentansprüche für folgenden Vertragsstaat : ES

 Verfahren zur Herstellung eines Fusionsproteins, das einen rekombinanten Einketten-Antik\u00f3rper umfa\u00dft, der gegen die extrazellul\u00e4re Dom\u00e4ne des Wachstumsfaktor-Rezeptors c-erbB-2 gerichtet ist, welcher eine variable Do-

mäne einer schweren Kette und eine variable Domäne einer leichten Kette eines monoklonalen Antikörpers, wobei die Domänen durch einen Polypeptid-Spacer verbunden sind, und eine Effektorgruppe umfaßt, dadurch gekennzeichnet, daß die Zellen, die einen derartigen Antikörper synthetisieren in vitro oder in vivo vermehrt werden und, wenn erforderlich, der erhaltene Antikörper isoliert wird.

2. Verfahren zur Herstellung eines Fusionsproteins nach Anspruch 1, worin die variable Dom\u00e4ne der schweren Kette ein Polypeptid der Aminos\u00e4uresequenz 2-120 der SEQ ID Nr.4 umfa\u00d6t, worin gegebenenfalls 1, 2, 3 oder 4 einzelne Aminos\u00e4uren in den Aminos\u00e4uresequenzen 2-31 (FR₁), 37-50 (FR₂), 68-99 (FR₃) und/ oder 110-120 (FR₄), durch andere Aminos\u00e4uren ersetzt sind, oder entfernt wurden, und worin die Aminos\u00e4ure Cys in dem oxidierten Zustand unter Bildung von S-S-Br\u00fccken vorliegen kann.

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- Verfahren zur Herstellung eines Fusionsproteins nach Anspruch 1, worin die variable Dom\u00e4ne der schweren Kette ein Polypeptid der Aminos\u00e4uresequenz 2-120 der SEQ ID Nr. 4 umfa\u00d6t, worin die Aminos\u00e4ure Cys in dem oxidierten Zustand unter Bildung von S-S-Br\u00fccken vorliegen kann.
- 4. Verfahren zur Herstellung eines Fusionsproteins nach Anspruch 1, worin die variable Domäne der leichten Kette ein Polypeptid der Aminosäuresequenz 136-241 der SEQ ID Nr.4 umfaßt, worin gegebenenfalls 1, 2, 3 oder 4 einzelne Aminosäuren in den Aminosäuresequenzen 136-158 (FR₆), 170-184 (FR₇), 192-223 (FR₈) und/oder 233-241 (FR₉) durch andere Aminosäuren ersetzt sind, oder entfernt wurden, und worin die Aminosäure Cys in dem oxidierten Zustand unter Bildung von S-S-Brücken vorliegen kann.
- Verfahren zur Herstellung eines Fusionsproteins nach Anspruch 4, worin die variable Domäne der leichten Kette ein Polypeptid der Aminosäuresequenz 136-241 der SEQ ID Nr.4 umfaßt, worin die Aminosäure Cys in dem oxidierten Zustand unter Bildung von S-S-Brücken vorliegen kann.
- 6. Verfahren zur Herstellung eines Fusionsproteins nach Anspruch 1, worin die variable Domäne der schweren Kette ein Polypeptid der Aminosäuresequenz 2-121 der SEQ ID Nr.8 umfaßt, worin gegebenenfalls 1, 2, 3 oder 4 einzelne Aminosäuren in den Aminosäuresequenzen 2-31 (FR₁), 37-50 (FR₂), 68-99 (FR₃) und/ oder 111-121 (FR₄) durch andere Aminosäuren ersetzt sind, oder entfernt wurden und worin die Aminosäure Cys in dem oxidierten Zustand unter Bildung von S-S- Brücken vorliegen kann.
- 7. Verfahren zur Herstellung eines Fusionsproteins nach Anspruch 6, worin die variable Domäne der schweren Kette ein Polypeptid der Aminosäuresequenz 2-121 der SEQ ID Nr.8 umfaßt, worin die Aminosäure Cys in dem oxidierten Zustand unter Bildung von S-S- Brücken vorliegen kann.
- 8. Verfahren zur Herstellung eines Fusionsproteins nach Anspruch 6, worin die variable Domäne der leichten Kette ein Polypeptid der Aminosäuresequenz 137-241 der SEQ ID Nr.8 umfaßt, worin gegebenenfalls eine oder mehrere einzelne Aminosäuren in den Aminosäuresequenzen 137-159 (FR₆), 171-185 (FR₇), 193-224 (FR₈) und/oder 233-241 (FR₉) durch andere Aminosäuren ersetzt sind, oder entfernt wurden, und worin die Aminosäure Cys in dem oxidierten Zustand unter Bildung von S-S-Brücken vorliegen kann.
- Verfahren zur Herstellung eines Fusionsproteins nach Anspruch 8, worin die variable Domäne der teichten Kette ein Polypeptid der Aminosäuresequenz 137-241 der SEQ ID Nr.8 umfaßt, worin die Aminosäure Cys in dem oxidierten Zustand unter Bildung von S-S-Brücken vorliegen kann.
- Verfahren zur Herstellung eines Fusionsproteins nach Anspruch 1, worin das Effektormolekül ein Enzym oder eine biologisch aktive Variante davon ist.
- 11. Verfahren nach Anspruch 10, worin das Enzym alkalische Phosphatase oder eine biologisch aktive Variante davon
- 12. Verfahren nach Anspruch 10, worin das Effektormolekül ein Toxin oder eine biologisch aktive Variante davon ist.
- Verfahren nach Anspruch 12, worin das Effektormolekül Pseudomonas Exotoxin oder eine biologisch aktive Variante davon ist.
  - 14. Verfahren zur Herstellung eines Fusionsproteins nach Anspruch 1, worin die variable Dom\u00e4ne der schweren Kette und die variable Dom\u00e4ne der leichten Kette von einem monoklonalen Maus-Antik\u00f6rper abgeleitet ist, der ausge-

wählt ist aus der Gruppe bestehend aus FRP5, FSP16, FWP51 und FSP77, die die gemäß dem Budapester Vertrag am 21. November 1990 bei der European Collection of Animal Cell Cultures (ECACC) in Porton Down, Salisbury, GB, unter den Hinterlegungsnummern 90112115, 90112116, 90112117 bzw. 90112118 hinterlegt wurden.

5 15. Verfahren nach Anspruch 1, worin die variable Dom\u00e4ne der schweren Kette und die variable Dom\u00e4ne der leichten Kette von dem monoklonalen Maus-Antik\u00f6rper FRP5 abgeleitet sind.

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- 16. Verfahren nach Anspruch 1, worin die variable Dom\u00e4ne der schweren Kette und die variable Dom\u00e4ne der leichten Kette von dem monoklonalen Maus-Antik\u00f6rper FWP51 abgeleitet sind.
- 17. Verfahren nach Anspruch 1 zur Herstellung eines Fusionsproteins, das als Fv(FRP5)-phoA bezeichnet wird, welches ein Polypeptid der Aminosäuresequenz 2-690 der SEQ ID Nr.5 umfaßt.
- Verfahren nach Anspruch 1 zur Herstellung eines Fusionsproteins, das als Fv(FRP5)-ETA 1 bezeichnet wird, welches ein Polypeptid der Aminosäuresequenz 2-606 der SEQ ID Nr.10 umfaßt.
- Verfahren nach Anspruch 1 zur Herstellung eines Fusionsproteins, das als Fv(FWP51)-ETA bezeichnet wird, welches ein Polypeptid der Aminosäuresequenz 2-606 der SEQ ID Nr.II umfaßt.
- 20. Verfahren nach Anspruch 1, worin die Zellen, die das Fusionsprotein exprimieren, mit einer rekombinanten DNA-Sequenz transformiert sind, die das Fusionsprotein codiert.
  - 21. Verfahren nach Anspruch 20, worin die rekombinante DNA-Sequenz ein Insert umfaßt, das eine variable Domäne einer schweren Kette der Maus eines monoklonalen Antikörpers codiert, der ausgewählt ist aus der Gruppe bestehend aus den Antikörpern FRP5, FSP16, FSP77 und FWP51, die gemäß dem Budapester Vertrag am 21. November 1990 bei der European Collection of Animal Cell Cultures (ECACC) in Porton Down, Salisbury, GB, unter den Hinterlegungsnummern 90112115, 90112116, 90112117 bzw. 90112118 hinterlegt wurden, oder eine Aminosäuresequenz codiert, die zu der variablen Domäne der schweren Kette homolog ist.
- 22. Verfahren nach Anspruch 20, worin die rekombinante DNA-Sequenz ein Insert umfaßt, das eine variable Domäne einer leichten Kette der Maus eines monoklonalen Antikörpers codiert, der ausgewählt ist aus der Gruppe bestehend aus den Antikörpern FRP5, FSP16, FSP77 und FWP51, die gemäß dem Budapester Vertrag am 21. November 1990 bei der European Collection of Animal Cell Cultures (ECACC) in Porton Down, Salisbury, GB, unter den Hinterlegungsnummern 90112115, 90112116, 90112117 bzw. 90112118 hinterlegt wurden, oder eine Aminosäuresequenz codiert, die zu der variablen Domäne der leichten Kette homolog ist.
  - 23. Verfahren nach Anspruch 20, worin die rekombinante DNA-Sequenz in einem Hybridvektor enthalten ist, der weiter einen Ursprung der Replikation oder eine autonom replizierende Sequenz, einen oder mehrere dominante Markersequenzen, und gegebenenfalls Expressionskontrollsequenzen, Signalsequenzen und zusätzliche Restriktionsschnittstellen umfaßt.
  - Verfahren nach Anspruch 23, worin der Hybridvektor weiter einen Affenvirus-Promotor und den Enhancer der Maus Ig H- oder L-Kette umfaßt.
- 45 25. Verfahren zur Herstellung einer rekombinanten DNA-Sequenz nach Anspruch 20, welches die folgenden Schritte
  - a) Herstellen von Maus-DNA aus dem Genom einer geeigneten Hybridomzellinie und Auswählen der gewünschten DNA, die die variablen Domänen der schweren und/oder leichten Kette des Antikörpers mit der gewünschten Spezifität codiert,
  - b) Herstellen von DNA, die die gewünschte Signalsequenz codiert und/oder Herstellen von DNA, die ein Effektormolekül codiert,
  - c) Synthetisieren von DNA mittels chemischer Methoden, die die gewünschte Spacergruppe codiert,
  - d) Bilden rekombinanter Gene, die die rekombinanten Antikörper codieren, indern die DNA aus Schritt a) und gegebenenfalls b) und/oder c) in geeignete Hybridvektoren eingebaut werden,
  - e) Transferieren der enthaltenen Hybridvektoren in eine aufnehmende Wirtszelle oder Wiedergewinnen der DNA, die die rekombinanten Gene codiert und Transferieren der nicht-verknüpften DNA in eine aufnehmende Wirtszelle

- f) Auswählen und Züchten der transformierten Wirtszelle, und
- g) gegebenenfalls Isolieren der gewünschten DNA.
- 26. Verfahren zur Herstellung einer Wirtszelle, die mit einer rekombinanten DNA-Sequenz nach Anspruch 20 transformiert ist, wobei geeignete aufnehmende Zellen mit einem Hybridvektor nach Anspruch 23, einem Ursprung der Replikation oder einer autonom replizierenden Sequenz, einem oder mehreren dominanten Markersequenzen und gegebenenfalls Expressionskontrollsequenzen, Signalsequenzen und zusätzlichen Restriktionsschnittstellen transformiert werden, und die transformierten Zellen ausgewählt werden.

### Revendications

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## Revendications pour les Etats contractants suivants : AT, BE, CH, LI, DE, DK, FR, GB, GR, IT, LU, NL, PT, SE

- 1. Protéine de fusion comprenant un anticorps recombinant monocaténaire dirigé contre le domaine extracellulaire du récepteur c-erbB-2 de facteur de croissance, comprenant un domaine variable de chaîne lourde et un domaine variable de chaîne légère d'un anticorps monoclonal, lesquels domaines sont reliés par un groupe espaceur polypeptidique, et une molécule effectrice, et comprenant éventuellement un peptide facilitant la purification, un site de coupure et un espaceur peptidique.
- 2. Protéine de fusion selon la revendication 1, dans laquelle le domaine variable de chaîne lourde comprend un polypeptide ayant la séquence d'aminoacides 2 à 120 d'ID SEQ nº 4, éventuellement 1, 2, 3 ou 4 aminoacides individuels dans les séquences d'aminoacides 2 à 31 (FR₁), 37 à 50 (FR₂), 68 à 99 (FR₃) et/ou 110 à 120 (FR₄) étant remplacés par d'autres aminoacides ou supprimés, et l'aminoacide Cys pouvant être à l'état oxydé, en formant des ponts S-S.
- Protéine de fusion selon la revendication 1, dans laquelle le domaine variable de chaîne lourde comprend un polypeptide ayant la séquence d'aminoacides 2 à 120 d'ID SEQ n° 4, dans lequel l'aminoacide Cys peut être à l'état oxydé, en formant des ponts S-S.
- 4. Protéine de fusion selon la revendication 1, dans laquelle le domaine variable de chaîne légère comprend un polypeptide ayant la séquence d'aminoacides 136 à 241 d'ID SEQ n° 4, éventuellement 1, 2, 3 ou 4 aminoacides individuels dans les séquences d'aminoacides 136 à 158 (FR₈), 170 à 184 (FR₇), 192 à 223 (FR₈) et/ou 233 à 241 (FR₉) étant remplacés par d'autres aminoacides ou supprimés, et l'aminoacide Cys pouvant être à l'état oxydé, en formant des ponts S-S.
- 5. Protéine de fusion selon la revendication 4, dans laquelle le domaine variable de chaîne légère comprend un polypeptide ayant la séquence d'aminoacides 136 à 241 d'ID SEQ n° 4, dans lequel l'aminoacide Cys peut être à l'état oxydé, en formant des ponts S-S.
  - 6. Protéine de fusion selon la revendication 1, dans laquelle le domaine variable de chaîne lourde comprend un polypeptide ayant la séquence d'aminoacides 2 à 121 d'ID SEQ n° 8, éventuellement 1, 2, 3 ou 4 aminoacides individuels dans les séquences d'aminoacides 2 à 31 (FR₁), 37 à 50 (FR₂), 68 à 99 (FR₃) et/ou 111 à 121 (FR₄) étant remplacés par d'autres aminoacides ou supprimés, et l'aminoacide Cys pouvant être à l'état oxydé, en formant des ponts S-S.
  - 7. Protéine de fusion selon la revendication 6, dans laquelle le domaine variable de chaîne lourde comprend un polypeptide ayant la séquence d'aminoacides 2 à 121 d'ID SEQ nº 8, dans lequel l'aminoacide Cys peut être à l'état oxydé, en formant des ponts S-S.
  - 8. Protéine de fusion selon la revendication 6, dans laquelle le domaine variable de chaîne légère comprend un polypeptide ayant la séquence d'aminoacides 137 à 241 d'ID SEQ nº 8, éventuellement un ou plusieurs aminoacides individuels dans les séquences d'aminoacides 137 à 159 (FR₆), 171 à 185 (FR₇), 193 à 224 (FR₈) eVou 233 à 241 (FR₉) étant remplacés par d'autres aminoacides ou supprimés, et l'aminoacide Cys pouvant être à l'état oxydé, en formant des ponts S-S.
  - 9. Protéine de fusion selon la revendication 8, dans laquelle le domaine variable de chaîne légère comprend un

polypeptide ayant la séquence d'aminoacides 137 à 241 d'ID SEQ n° 8, dans lequel l'aminoacide Cys peut être à l'état coydé, en formant des ponts S-S.

10. Protéine de fusion selon l'une quelconque des revendications 1 à 9, dans laquelle la molécule effectrice est une enzyme ou un variant biologiquement actif de celle-ci.

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- Protéine de fusion selon la revendication 10, dans laquelle l'enzyme est la phosphatase alcaline ou un variant biologiquement actif de celle-ci.
- 10 12. Protéine de fusion selon l'une quelconque des revendications 1 à 9, dans laquelle la molécule effectrice est une toxine ou un variant biologiquement actif de celle-ci.
  - 13. Protéine de fusion selon la revendication 12, dans laquelle la molécule effectrice est l'exotoxine de Pseudomonas ou un variant biologiquement actif de celle-ci.
  - 14. Protéine de fusion selon la revendication 1, dans laquelle le domaine variable de chaîne lourde et le domaine variable de chaîne légère proviennent d'un anticorps monoclonal de souris choisi dans le groupe constitué par FRP5, FSP16, FWP51 et FSP77 déposés, sous les clauses du Traité de Budapest, le 21 novembre 1990 à l'European Collection of Animal Cell Cultures (ECACC), Porton Down, Salisbury, GB, sous les numéros de dépôt 90112115, 90112116, 90112117 et 90112118, respectivement.
  - 15. Protéine de fusion selon la revendication 1, dans laquelle le domaine variable de chaîne lourde et le domaine variable de chaîne légère proviennent de l'anticorps monoclonal FRP5 de souris.
- 25 16. Protéine de fusion selon la revendication 1, dans laquelle le domaine variable de chaîne lourde et le domaine variable de chaîne légère proviennent de l'anticorps monoclonal FWP51 de souris.
  - Protéine de fusion désignée par Fv(FRP5)-phoA selon la revendication 1, comprenant un polypeptide ayant la séquence d'aminoacides 2 à 690 d'ID SEQ n° 5.
  - 18. Protéine de fusion désignée par Fv(FRP5)-ETA selon la revendication 1, comprenant un polypeptide ayant la séquence d'aminoacides 2 à 606 d'ID SEQ n° 10.
  - 19. Protéine de fusion désignée par Fv(FWP51)-ETA selon la revendication 1, comprenant un polypeptide ayant la séquence d'aminoacides 2 à 606 d'ID SEQ n° 11.
  - 20. ADN recombinant comprenant un insert codant pour une protéine de fusion selon l'une quelconque des revendications 1 à 19.
- 21. ADN recombinant selon la revendication 20, comprenant un insert codant pour un domaine variable murin de chaîne lourde d'un anticorps monoclonal choisi dans le groupe constitué par les anticorps FRP5, FSP16, FSP77 et FWP51 déposés, sous les clauses du Traité de Budapest, le 21 novembre 1990 à l'European Collection of Animal Cell Cultures (ECACC), Porton Down, Salisbury, GB, sous les numéros de dépôt 90112115, 90112116, 90112117 et 90112118, respectivement, ou codant pour une séquence d'aminoacides homologue dudit domaine variable de chaîne lourde.
  - 22. ADN recombinant selon la revendication 20, comprenant un insert codant pour un domaine variable murin de chaîne légère d'un anticorps monoclonal choisi dans le groupe constitué par les anticorps FRP5, FSP16, FSP77 et FWP51 déposés, sous les clauses du Traité de Budapest, le 21 novembre 1990 à l'European Collection of Animal Cell Cultures (ECACC), Porton Down, Salisbury, GB, sous les numéros de dépôt 90112115, 90112116, 90112117 et 90112118, respectivement, ou codant pour une séquence d'aminoacides homologue dudit domaine variable de chaîne légère.
- 23. ADN recombinant selon la revendication 20, qui est un vecteur hybride comprenant en outre une origine de réplication ou une séquence à réplication autonome, une ou plusieurs séquences de marquage dominantes et, éventuellement, des séquences régulatrices d'expression, des séquences signal et des sites de restriction supplémentaires.

- 24. Vecteur hybride selon la revendication 23, comprenant un promoteur de virus simien et l'activateur de chaîne lourde ou de chaîne légère d'Ig de souris.
- 25. Procédé pour la production d'un ADN selon la revendication 20, comprenant les étapes suivantes:
  - a) production d'ADN murin à partir du génome d'une lignée d'hybridomes appropriée, et sélection de l'ADN recherché codant pour les domaines variables de chaîne légère et/ou de chaîne lourde de l'anticorps ayant la spécificité désirée,
  - b) production d'ADN codant pour la séquence signal désirée et production d'ADN codant pour une molécule effectrice,
  - c) synthèse d'ADN codant pour le groupe espaceur désiré, par des méthodes chimiques,
  - d) construction de gènes recombinants codant pour les protéines de fusion, par incorporation de l'ADN des étapes a) et c) et éventuellement b), dans des vecteurs hybrides appropriés,
  - e) transfert des vecteurs hybrides obtenus, dans une cellule hôte réceptrice, ou récupération de l'ADN codant pour les gènes recombinants et transfert de l'ADN non lié, dans une cellule hôte réceptrice,
  - f) sélection et culture de la cellule hôte transformée, et
  - g) éventuellement isolement de l'ADN recherché.

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- 26. Cellule hôte transformée par un ADN recombinant selon la revendication 25.
- 27. Cellule hôte selon la revendication 26, qui est une cellule d'une souche de E. coli.
- 28. Procédé pour la production d'une cellule hôte transformée, selon la revendication 6, dans lequel des cellules réceptrices appropriées sont transformées par un vecteur hybride comprenant un insert d'ADN selon la revendication 20, une origine de réplication ou une séquence se répliquant de façon autonome, une ou plusieurs séquences de marquage dominantes et, éventuellement, des séquences régulatrices d'expression, des séquences signal et des sites de restriction supplémentaires, et les cellules transformées sont sélectionnées.
- 29. Utilisation d'une protéine de fusion selon la revendication 1, pour la détermination qualitative et quantitative du récepteur c-erbB-2 de facteur de croissance.
  - 30. Utilisation selon la revendication 29, comprenant la coloration immunologique de coupes de tissu, à l'aide d'une solution contenant la protéine de fusion comprenant une enzyme détectable.
- 35 31. Nécessaire d'essai pour la détermination qualitative et quantitative de la protéine c-erbB-2, comprenant une protéine de fusion selon la revendication 1.
  - 32. Protéine de fusion selon la revendication 1, pour utilisation pour le traitement de l'organisme humain ou animal.
- 33. Composition pharmaceutique pour le traitement de tumeurs exprimant excessivement le récepteur c-erbB-2 de facteur de croissance, comprenant une quantité thérapeutiquement efficace d'une protéine de fusion selon la revendication 1, et un véhicule pharmaceutiquement acceptable.
  - 34. Utilisation d'une protéine de fusion selon la revendication 1, pour la préparation d'une composition pharmaceutique.

### Revendications pour l'Etat contractant suivant : ES

- 1. Procédé pour la production d'une protéine de fusion comprenant un anticorps recombinant monocaténaire dirigé contre le domaine extracellulaire du récepteur c-erbB-2 de facteur de croissance, comprenant un domaine variable de chaîne lourde et un domaine variable de chaîne légère d'un anticorps monoclonal, qui sont reliés par un espaceur polypeptidique, et un groupe effecteur, caractérisé en ce que des cellules productrices d'un tel anticorps sont multipliées in vitro ou in vivo et, si nécessaire, l'anticorps obtenu est isolé.
- 2. Procédé selon la revendication 1, pour la production d'une protéine de fusion dans laquelle le domaine variable de chaîne lourde comprend un polypeptide ayant la séquence d'aminoacides 2 à 120 d'ID SEQ nº 4, éventuellement 1, 2, 3 ou 4 aminoacides individuels dans les séquences d'aminoacides 2 à 31 (FR₁), 37 à 50 (FR₂), 68 à 99 (FR₃) et/ou 110 à 120 (FR₄) étant remplacés par d'autres aminoacides ou supprimés, et l'aminoacide Cys

pouvant être à l'état oxydé, en formant des ponts S-S.

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- 3. Procédé selon la revendication 1, pour la production d'une protéine de fusion laquelle le domaine variable de chaîne lourde comprend un polypeptide ayant la séquence d'aminoacides 2 à 120 d'ID SEQ nº 4, dans lequel l'aminoacide Cys peut être à l'état oxydé, en formant des ponts S-S.
- 4. Procédé selon la revendication 1, pour la production d'une protéine de fusion dans laquelle le domaine variable de chaîne légère comprend un polypeptide ayant la séquence d'aminoacides 136 à 241 d'ID SEQ n° 4, éventuellement 1, 2, 3 ou 4 aminoacides individuels dans les séquences d'aminoacides 136 à 158 (FR₆), 170 à 184 (FR₇), 192 à 223 (FR₈) et/ou 233 à 241 (FR₉) étant remplacés par d'autres aminoacides ou supprimés, et l'aminoacide Cys pouvant être à l'état oxydé, en formant des ponts S-S.
- 5. Procédé selon la revendication 4, pour la production d'une protéine de fusion dans laquelle le domaine variable de chaîne légère comprend un polypeptide ayant la séquence d'aminoacides 136 à 241 d'ID SEQ n° 4, dans lequel l'aminoacide Cys peut être à l'état oxydé, en formant des ponts S-S.
- 6. Procédé selon la revendication 1, pour la production d'une protéine de fusion dans laquelle le domaine variable de chaîne lourde comprend un polypeptide ayant la séquence d'aminoacides 2 à 121 d'ID SEQ nº 8, éventuellement 1, 2, 3 ou 4 aminoacides individuels dans les séquences d'aminoacides 2 à 31 (FR₁), 37 à 50 (FR₂), 68 à 99 (FR₃) eVou 111 à 121 (FR₄) étant remplacés par d'autres aminoacides ou supprimés, et l'aminoacide Cys pouvant être à l'état oxydé, en formant des ponts S-S.
- 7. Procédé selon la revendication 6, pour la production d'une protéine de fusion dans laquelle le domaine variable de chaîne lourde comprend un polypeptide ayant la séquence d'aminoacides 2 à 121 d'ID SEQ nº 8, dans lequel l'aminoacide Cys peut être à l'état oxydé, en formant des ponts S-S.
- 8. Procédé selon la revendication 6, pour la production d'une protéine de fusion dans laquelle le domaine variable de chaîne légère comprend un polypeptide ayant la séquence d'aminoacides 137 à 241 d'ID SEQ n° 8, éventuellement un ou plusieurs aminoacides individuels dans les séquences d'aminoacides 137 à 159 (FR₆), 171 à 185 (FR₇), 193 à 224 (FR₈) et/ou 233 à 241 (FR₉) étant remplacés par d'autres aminoacides ou supprimés, et l'aminoacide Cys pouvant être à l'état oxydé, en formant des ponts S-S.
- Procédé selon la revendication 8, pour la production d'une protéine de fusion dans laquelle le domaine variable de chaîne légère comprend un polypeptide ayant la séquence d'aminoacides 137 à 241 d'ID SEQ n° 8, dans lequel l'aminoacide Cys peut être à l'état oxydé, en formant des ponts S-S.
- 10. Procédé selon la revendication 1, pour la production d'une protéine de fusion dans laquelle la molécule effectrice est une enzyme ou un variant biologiquement actif de celle-ci.
- Procédé selon la revendication 10, dans lequel l'enzyme est la phosphatase alcaline ou un variant biologiquement actif de celle-ci.
  - Procédé selon la revendications 10, dans lequel la molécule effectrice est une toxine ou un variant biologiquement actif de celle-ci.
  - Procédé selon la revendication 12, dans lequel la molécule effectrice est l'exotoxine de Pseudomonas ou un variant biologiquement actif de celle-ci.
  - 14. Procédé selon la revendication 1, pour la production d'une protéine de fusion dans laquelle le domaine variable de chaîne lourde et le domaine variable de chaîne légère proviennent d'un anticorps monoclonal de souris choisi dans le groupe constitué par FRP5, FSP16, FWP51 et FSP77 déposés, sous les clauses du Traité de Budapest, le 21 novembre 1990 à l'European Collection of Animal Cell Cultures (ECACC), Porton Down, Salisbury, GB, sous les numéros de dépôt 90112115, 90112116, 90112117 et 90112118, respectivement.
  - 15. Procédé selon la revendication 1, dans lequel le domaine variable de chaîne lourde et le domaine variable de chaîne légère proviennent de l'anticorps monoclonal FRP5 de souris.
    - 16. Procécé selon la revendication 1, dans lequel le domaine variable de chaîne lourde et le domaine variable de

chaîne légère proviennent de l'anticorps monoclonal FWP51 de souris.

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- 17. Procédé selon la revendication 1, pour la production d'une protéine de fusion désignée par Fv(FRP5)-phoA, comprenant un polypeptide ayant la séquence d'aminoacides 2 à 690 d'ID SEQ n° 5.
- 18. Procédé selon la revendication 1, pour la production d'une protéine de fusion désignée par Fv(FRP5)-ETA c mprenant un polypeptide ayant la séquence d'aminoacides 2 à 606 d'ID SEQ n° 10.
- Procédé selon la revendication 1, pour la production d'une protéine de fusion désignée par Fv(FWP51)-ETA comprenant un polypeptide ayant la séquence d'aminoacides 2 à 606 d'ID SEQ n° 11.
  - 20. Procédé selon la revendication 1, dans lequel les cellules exprimant la protéine de fusion sont transformées par une séquence d'ADN recombinant codant pour ladite protéine de fusion.
- 21. Procédé selon la revendication 20, dans lequel la séquence d'ADN recombinant comprend un insert codant pour un domaine variable murin de chaîne lourde d'un anticorps monoclonal choisi dans le groupe constitué par les anticorps FRP5, FSP16, FSP77 et FWP51 déposés, sous les clauses du Traité de Budapest, le 21 novembre 1990 à l'European Collection of Animal Cell Cultures (ECACC), Porton Down, Salisbury, GB, sous les numéros de dépôt 90112115, 90112116, 90112117 et 90112118, respectivement, ou codant pour une séquence d'aminoacides homologue dudit domaine variable de chaîne lourde.
  - 22. Procédé selon la revendication 20, dans lequel la séquence d'ADN recombinant comprend un insert codant pour un domaine variable murin de chaîne légère d'un anticorps monoclonal choisi dans le groupe constitué par les anticorps FRP5, FSP16, FSP77 et FWP51 déposés, sous les clauses du Traité de Budapest, le 21 novembre 1990 à l'European Collection of Animal Cell Cultures (ECACC), Porton Down, Salisbury, GB, sous les numéros de dépôt 90112115, 90112116, 90112117 et 90112118, respectivement, ou codant pour une séquence d'aminoacides homologue dudit domaine variable de chaîne légère.
  - 23. Procédé selon la revendication 20, dans lequel la séquence d'ADN recombinant est comprise dans un vecteur hybride comprenant en outre une origine de réplication ou une séquence à réplication autonome, une ou plusieurs séquences de marquage dominantes et, éventuellement, des séquences régulatrices d'expression, des séquences signal et des sites de restriction supplémentaires.
    - 24. Procédé selon la revendication 23, dans lequel le vecteur hybride comprend en outre un promoteur de virus simien et l'activateur de chaîne lourde ou de chaîne légère d'Ig de souris.
  - 25. Procédé pour la production d'une séquence d'ADN recombinant selon la revendication 20, comprenant les étapes suivantes:
    - a) production d'ADN murin à partir du génome d'une lignée d'hybridomes appropriée, et sélection de l'ADN recherché codant pour les domaines variables de chaîne légère el/ou de chaîne lourde de l'anticorps ayant la spécificité désirée.
      - b) production d'ADN codant pour la séquence signal désirée et/ou production d'ADN codant pour une molécule effectrice
      - c) synthèse d'ADN codant pour le groupe espaceur désiré, par des méthodes chimiques,
      - d) construction de gènes recombinants codant pour les anticorps recombinants, par incorporation de l'ADN de l'étape a) et éventuellement b) et/ou c), dans des vecteurs hybrides appropriés,
      - e) transfert des vecteurs hybrides obtenus, dans une cellule hôte réceptrice, ou récupération de l'ADN codant pour les gènes recombinants et transfert de l'ADN non lié, dans une cellule hôte réceptrice,
      - f) sélection et culture de la cellule hôte transformée, et
      - g) éventuellement isolement de l'ADN recherché.
  - 26. Procédé pour la production d'une cellule hôte transformée par une séquence d'ADN recombinant selon la revendication 20, dans lequel des cellules réceptrices appropriées sont transformées par un vecteur hybride selon la revendication 23, une origine de réplication ou une séquence se répliquant de façon autonome, une ou plusieurs séquences de marquage dominantes et, éventuellement, des séquences régulatrices d'expression, des séquences signal et des sites de restriction supplémentaires, et les cellules transformées sont sélectionnées.